REMOTE SENSING OF FORESTS

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PREFACE

This compendium has been generously financed by the Erik Johan Ljungbergs Educational fund. It has been part of a larger effort to introduce more technical competence into the Swedish University of Agricultural Sciences (SLU) Forestry Master’s program. The contributors to the compendium come from the Department of Forest Resource Management at SLU in Umeå. The compendium consists of three main parts: Forest Inventory, Remote Sensing of Forests, and Forest Planning. It is intended to be used as literature in the Department’s courses, primarily at the basic undergraduate level, but is also freely available on the internet and can be used by general readers interested in the topics. The compendium should give the reader an understanding of basic concepts in the three areas individually, and as a whole, and give insight into the process of creating forest information that could be used in assessment and planning of forest resources, including input to Heureka or GIS.

In the remote sensing part of the compendium, we aim to answer the following questions: How can forest data be effectively yet accurately collected? What data do forest industry or government authorities need? What are advantages or pitfalls of different data and methods? Sometimes a sample of the landscape via a field-based inventory is all that is needed. In other cases, full area coverage data are needed. Remote sensing (RS) can provide full area coverage on, for example, tree height, location of harvested stands, stand boundaries, forest health, tree species, and forest structure. The current trend in RS is 3D analysis with laser scanning, digital photogrammetry, or radar data, while vegetation types can be seen using optical satellite data. To process RS data, inventory data are needed to link ground data with what can be seen from the air. On the other hand, samples can be taken from RS data to produce statistics. Advances in sensor-based field inventory are being made with terrestrial laser scanning. What do these data deliver compared to field-based inventory?

The remote sensing compendium will first address background concepts and history of remote sensing, as well as remote sensing’s relationship to forest inventory and forest planning in Chapter 1. The physics of remote sensing is covered in Chapter 2. Basic information regarding sensors, platforms and digital data are discussed in Chapter 3. Chapter 4 goes into more detail about electro-optical sensors, and Chapter 5 discusses traditional aerial photography interpretation, as well as digital manipulation into three-dimensional point data. Chapter 6 takes up airborne and terrestrial laser scanning, while Chapter 7 addresses radar data. Chapter 8 focusses on issues of combining reference data and remote sensing data, and desirable properties of the reference data. Chapter 9 gives details on the methods used to combine reference data and remote sensing data in order to process the data and produce new information. Chapter 10 discusses the methods and importance of accuracy assessment of the information produced. Chapter 11 gives a short introduction to programming used currently in remote sensing.
The remote sensing part of the compendium also will use a framework we see as helpful in referring to remote sensing image processing. That framework is given below and will be referred to in the text.

We hope that this compendium informs the reader about the past and current knowledge, as well as the future potential, of Forest Remote Sensing.

Umeå, Sweden
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Heather Reese
Senior Lecturer in Forest Remote Sensing

Håkan Olsson
Professor in Forest Remote Sensing
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1. INTRODUCTION TO REMOTE SENSING

**Remote sensing.** The term used to describe the act of observing a phenomenon from a distance. Remote sensing has often dealt with the imaging of areas on the Earth’s surface taken by sensors placed on airplanes or satellites; however, it is becoming all the more common with sensors placed on moving vehicles or on ground-based platforms.

**Electromagnetic spectrum.** The entire range of wavelengths or frequencies of electromagnetic radiation extending from gamma rays to the longest radio waves and including visible light.

**Sensor.** The instrument which detects and stores observations.

**Platform.** The carrier of the sensor. This may be, for example, a stative, a small airplane, or a satellite.

1.1. What is remote sensing?

Remote sensing is the name of the subject which deals with observing objects remotely. Remote sensing is used in many applications including forestry, weather, nature conservation, cartography, just to name a few. Remote sensing is often performed via a sensor which observes and records properties of the electromagnetic spectrum. The sensor is placed on a platform, which can be, for example, a satellite or an airplane. The information recorded from the sensor is then analyzed in order to produce information usable as maps or within GIS or planning systems.

There are several definitions of remote sensing found in different textbooks, each with a little different wording. The following two definitions are similar and rather good:

> "**Remote sensing is the science and art of obtaining information about an object, area, or phenomenon through the analysis of data acquired by a device that is not in contact with the object, area or phenomenon under investigation**."  

This in fact leaves a rather wide-open definition. It can include many different aspects of remote sensing such as walking through a forest looking with your eyes, taking photographs with your pocket camera, or acquiring data with laser sensors flown in a plane or from a satellite flying hundreds of kilometers above Earth in space.

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In Swedish, the word for remote sensing is “fjärranalys”, which can be broken down into two English words, “fjärr = remote” and “analys = analysis”. In Lillesand et al.’s definition we find the important word “analysis”. While some definitions define remote sensing as an act which collects data, the definition can be widened to include the act of analyzing data in order to obtain useful information.

As a rule, this analysis is done by computer-based methods. For example, in a first step special software can be used to calculate the characteristics collected from remote sensing data, such as the ratio between the values of two wavelengths recorded within each pixel (e.g., red value/near-infrared value), or the difference between the highest and lowest laser return within an area. Within forest remote sensing, it is usual that the information from the remote sensing data is translated into forest measurements with the support of field reference data, so that certain combinations of colors are translated to forest species classes, or the distance between the laser returns translated, for example, into basal mean height. Often, conventional statistical software, such as Minitab or R can be used for calibration of the remote sensing data against field data.

1.2. Why use remote sensing?

We can also ask the question “Why?” about remote sensing. Why should we use remote sensing? Often there must be some real benefit to using these data, which are not exactly simple to collect or analyze. Some of the benefits of remote sensing are that they can provide:

- **Synoptic views** This is a big word to explain that we get a kind of “bird’s eye view” over an area. A dictionary definition of “synoptic” is “giving a general view of a whole -- or -- displaying conditions as they exist simultaneously over a broad area”. Depending on the users’ needs and the sensor’s capabilities, the remote sensing data may cover very large areas, or be concentrated on a smaller area.

- **A cost-effective way to get information.** This is often an argument for use of remotely sensed data, as it can mean less field work than otherwise required and therefore less costs. However, note that field work IS still often needed for work with remote sensing (it’s not obsolete).

- **New information.** The Infrared bands provide a great deal of information about vegetation. Remote sensing data that observes these wavelengths can give new information we would not otherwise see with our own eyes. In addition, laser data give us highly accurate height measurements – in many cases more accurate than can be manually measured.

- **Coverage of remote areas.** When the sensor is mounted on a plane or satellite, for example, images can be
obtained regardless of whether there are roads or footpaths to travel to that area.

- **Objectivity.** Data are collected without the initial interpretation by human judgment, and provide a raw, objective source of digital data. Analyses – such as vegetation classification or estimation of wood volume - can then be applied to the digital data in an objective manner. Digital data lend themselves to automated computer analysis.

- **Repeated images.** As long as the same sensor and same settings are used, images over a time period can be used to look at a landscape over time, which can be considered as another dimension (the temporal dimension).

Cost efficiency is likely the most important reason to use remote sensing in forestry primarily because the need for manual measurements in the field can be reduced, even if the field references are also needed for remote sensing estimates. A prerequisite for the acceptance of remote sensing in practical forestry has, however, proved to be that it has been cost effective and provide at least as good data as provided by traditional inventory methods -- this is the case for the estimation of several forest variables using laser scanner data.

Assuming that the proposed collection method is cost effective, it is of course also beneficial if you can get even better data than with traditional methods. Most remote sensing methods provide data with better spatial resolution than traditional forest maps, which can give, for example, indications that only parts of a stand need to be thinned, or are damaged. In the case of estimation of tree height or wood volume with laser scanner data, one can also expect better estimation accuracy than with subjective manual methods. It is also an advantage that remote sensing estimates can be impartial and independent of the person who is carrying out the measurements.

In particular, the satellite based remote sensing covers large areas regularly. Most remote sensing satellites systematically cover all the Earth with a two to three week interval. Satellite-based remote sensing is particularly important for nationwide or international applications.

A regular supply of digital data that can be processed in computers also makes it quite easy to automatically detect areas that have changed in an unusual way, e.g., areas of damaged forest. However as a rule additional information (e.g., field or reference data) is required to determine the cause and exact nature of the detected changes.
1.3. History

In the late 1800s people attempted taking photographs from hot air balloons. Aerial photography technology was then developed rapidly during the First World War (WWI). After WWI aerial photographs began to be used for forest mapping in e.g., Canada in the 1920s. In Sweden aerial photography was tested for the first time in 1930 and even then, the application for forest use was central. However, it would take until after World War II before aerial photography use for forestry would really take off. In the 1950s, a government committee was formed that would promote the use of aerial photographs for forestry. Even today, the manual interpretation of aerial photographs is the dominant use of remote sensing in forestry. Mainly orthophotos are used, but the interpretation can also be made with different types of stereo-viewing instruments.

In the 1980s, Sweden launched the so-called LMV-method of forest variable estimation. It means that the forest is divided into compartments, the tree height is measured, and the estimate of tree density and species mix is done manually on a photogrammetric workstation. From these data, the timber volume is then estimated. The method was used among other things as SCA and Holmen redid the division of their holdings in the 1990s. Since the early 2000s, aerial photography has been acquired via digital cameras, which among other things has resulted in increased access to aerial photographs in color infrared.

Since the mid-00s the use of drones (also called UAVs or UAS) for aerial photography has increased. Contributing to the development of UAV remote sensing is a series of leaps in technology: the development of improved battery technology that permits the use of electric motors, as well as small navigation systems and digital cameras.

The first military remote sensing satellites came in 1960, and in 1972 Landsat 1 came which was the first civil earth resource satellite. Landsat 1 had pixels with an 80 x 80 m size. This spatial resolution was improved by the Landsat TM sensor which came in 1982 and had 30 x 30 m pixels, and SPOT 1 which was launched in 1986 and had 10 x 10 m pixels. These older satellite images are archived, and this archive can be very useful for the study of landscape development over time. During the early 2000’s, Landsat data became available free of cost, which has increased their use.

In the 1990s, satellite imagery with 1 x 1m pixels became available, however, these satellites were frequently operated by commercial actors (the data cost money), and have covered generally smaller areas than the Earth resource satellites such as Landsat.

Satellites are taking frequent pictures of the Earth, but a limitation of photographic-like images (i.e., color or color infrared imaging) from space
is that the images are often cloudy. There is therefore great interest in radar satellites, which can record data through clouds. The testing of different types of radar data for forest mapping has been going on since the 1990s; while the potential is great, the practical use of this technology is still limited.

In the mid-1990s airborne laser scanning also developed. This technique involves laser beams sent out from an instrument (placed within an airborne platform such as a plane or helicopter), where the distance is measured from the source of the laser beam down to the ground or canopy (or whatever object that laser beam is reflected from). The result is a three-dimensional (3D) point cloud. The measurement is very accurate and an important factor for the airborne laser technology has been developed is that although the aircraft's position and movement in the air is calculated using GPS and inertial navigation. Measuring forest with airborne laser was first tested in Leningrad in the late 1970s and in the United States and Canada in the mid-1980s. They used the instrument to just measure a profile from the aircraft (i.e, a profiling laser). In Sweden we were the first in 1991 to test an experimental system with a scanning laser, but it was not until commercial systems with integrated positioning came in the mid-1990s as the development of laser data use for forestry (and many other applications) took off.

Laser scanning first began to be used as an operational method for forest inventory in Norway in 2002. The method then took off with the newly introduced so-called “area-based approach”, which means that field measured reference plots are used to assign forest data to a combination of metrics calculated from the laser point cloud. In Sweden the first attempt was made on a larger scale by the Forest Agency and the SLU (10,000 ha for re-parcelling in Dalarna) in 2003. This was followed by a period when several forest companies tested the laser scanning in areas of approximately 10,000 ha. When the National Land Survey laser scanning started in 2009 forestry companies then had new access to cheap laser data, which led to several forestry companies ordering laser data (or the finished product of estimates) for their entire holding. First was Bergvik, who in 2011 decided to make airborne laser data estimates for their entire forest holding.

1.4. Current role of remote sensing to provide forest information

Aerial photographs, primarily digital orthophotos, are the main source of information for forest data. Orthophotos can be used for manual classification of compartments and as a background for a variety of digital map products. To some extent, manual measurement and interpretation of stereo aerial photographs in digital photogrammetric workstation is still performed according to the so-called LMV-method. Forest data estimation using this method, however, requires great skill of the photo-interpreter if it is to replace field work. An alternative is to use aerial photos or orthophotos for a preliminary evaluation of the area before field work.
Digital stereo aerial photographs can also be processed automatically by the computer to create a 3D point cloud similar to that obtained by airborne laser data. The Swedish National Land Survey starts in 2016 to produce such 3D point clouds from their aerial photography product which they will call a "surface model from aerial photographs." They are also referred to as 3D data from image matching. These data can be used to calculate a raster with tree canopy heights. They can also be used for forestry estimates in the same way as 3D point cloud from laser data. However, the data obtained from aerial photography does not give as good information about forest density as laser data does, and therefore, estimates of timber volume will be worse from 3D image-matching data.

Drones (Unmanned Aerial Vehicles or UAVs) is being put into use in the preparation of forest management plans for individual properties. It is so far mainly orthophotos created from mosaicking drone flyovers that are being used.

The availability of frequent and free satellite images has increased. In 2014 Landsat 8 was launched and in 2015 the European Sentinel 2A satellite with 10 m pixels was launched. Since about 2000 Skogsstyrelsen (The Swedish Forest Agency) has used satellite images to annually map all new clearcuts. The Swedish Forest Agency also uses satellite images to estimate the forest cleaning/thinning needed.

For the years 2000, 2005 and 2010 SLU has made national forest maps with 25 x 25 m grid cells by joint processing of satellite images and reference plots from the National Forest Inventory (NFI). These forest maps have been widely used among researchers and authorities, but estimates of forest data based on the use of "2D" optical satellite images from a time are generally not accurate enough to be used in forestry planning.

Lantmäteriet's (The National Mapping Agency) laser scanning throughout Sweden that started in 2009 will in practice be finished for the forested part of the country in 2016. What remains is a few areas in the mountains. This laser data can be purchased for a marginal cost by commercial actors and is free to governmental authorities. The national laser scanning has thus been a catalyst for the use of laser data in Swedish forestry. Bergvik and Holmen have ordered estimates for their entire holdings and SCA has ordered laser estimates based on National Land Survey data for their forests in Norrland. Together, the Forest Agency and SLU worked in the context of a government commission to make forest estimates for the country, based on Lantmäteriets laser scanner data and reference plots from the National Forest Inventory. Together with other consulting companies, the Forest Agency has also developed raster layers from laser data showing soil wetness maps, as well as a raster layer with tree heights at a 2 x 2 m grid cell size. These map layers can be viewed and downloaded from the Forestry Board's website. They have been widely used, both in the forestry sector and by other actors, including banks and insurance companies.
Currently there is much ongoing research and testing on a range of technologies with great potential to streamline the future measurement of forest. An example of this is that laser scanners can also be placed on a tripod on the ground (the name of this technology is often shortened to TLS for Terrestrial Laser Scanning), or mounted in a backpack or on a vehicle (often abbreviated to MLS for Mobile Laser Scanning), or even carried in the hand (also MLS). With TLS and MLS detailed information can be obtained about tree stem position and shape, which, inter alia, can be used as reference data to create estimates from the airborne data.

1.6. Products from remote sensing

Remote sensing provides raw data which can be used simply as a visual background image, such as it is done now with Google Earth. However, remote sensing data are digital data, and can be analysed and manipulated and therefore converted into map data that give users the information they need.

We can categorize map data as being thematic or continuous. Thematic maps have discrete classes, such as the Swedish Land Cover map with the thematic classes of “forest”, “mire”, “water”, for example. Continuous variable maps consist of a range of values for a single phenomenon. An example of this is a map of timber volume, with values ranging from 0 up to the maximum value. These two map outputs are demonstrated in Fig. 1.1.

Another common product from remote sensing data is a change map, which can show the differences between two or more dates of remote sensing data. Another product from remote sensing data is as input to visualization; the remote sensing data input may be as raw data, such as from Terrestrial Laser Scanning, or it may be the map products that form the baseline for visualization.

Fig 1.1. Example of a discrete or thematic value map on left (e.g., land cover classes), and a continuous value map on right (e.g., total wood volume).
1.7. The relationship between remote sensing and GIS, forest inventory and forest planning

The field of remote sensing has a relationship with other fields, such as Geographical Information Systems/Technology (GIS/GIT), forest inventory, and forest planning. It is worth clarifying the definition and roll of remote sensing in relation to these other fields.

A GIS consists of four components, namely data acquisition, data storage, data analysis, and map production\(^2\). Remote sensing fills the function of “data acquisition” in a GIS; the remote sensing data input may be raw data (e.g., images) or processed data (e.g., map data derived from remote sensing).

The field of forest inventory is concerned with techniques and methods for measuring and estimation of forest resources. Forest inventory can be done with manual methods, but remote sensing plays an increasing role for providing both wall-to-wall data and as ancillary data in statistical estimations of forest resources. Remote sensing and forest inventory have an intertwined relationship, which can be described as having two main interactions:

- Remote sensing data acquisition for forest inventory purposes, and
- Forest inventory data use as reference data to help interpret remote sensing data (i.e., training data) or to use in validation of the map or product from remote sensing (i.e., validation data).

Note that the subject area of remote sensing differs from the subject areas of inventory, planning, and GIS. Remote sensing data are used within the process of forest inventory, and they are analysed or displayed within a GIS. However, the subject of remote sensing includes not just measuring (i.e., inventory), but also knowledge of how to acquire and process the raw remotely sensed in a correct way. This may require background knowledge in physics, statistics, photogrammetry, programming, and certainly geography. The subject of remote sensing also involves knowing which remote sensing data source is best suited for the purpose (i.e., strengths and limitations), and how to perform and present an accuracy assessment of the map products from remote sensing data.

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1.8. Remote sensing for global to individual tree applications

The remote sensing data source chosen is dependent upon the aim and goal of the project, the availability of remote sensing data, and the cost, among other factors. For example, if we consider remote sensing data acquisition for forest inventory purposes, we need to consider at what scale we wish to produce information. This can be at the landscape scale and may range downwards to the individual tree scale.

The large area, landscape scale coverage that can be provided by remote sensing (synoptic views) makes it a useful tool. Aerial photographs have been used between 1930’s to present for delineating forest stands and measurement of tree height. Satellite data have also played a role in providing forest information over estates, or whole countries. With the current innovation of airborne LiDAR, which measures tree heights and forest density with high accuracy, the use of remote sensing for forest inventory is increasing rapidly. The availability of data to both private forest owners and larger companies means that they are used by both.

At the individual tree scale, remote sensing technologies are providing data from both the air and from the ground. The ground-based remote sensing includes terrestrial laser scanning, and ground-based photogrammetry. These sensors may be placed on platform which is a stative, or may be mobile (e.g., placed on a car or hand-held). The level of spatial detail in the remote sensing data, the accuracy of the map products, and cost-effectiveness have had an influence on whether the data will be useful for forestry applications.

1.9. The growth of remote sensing for forestry purposes

Developments in the subject and use of remote sensing is growing exponentially due to:

- the ability to acquire highly accurate and useful 3D data (from laser, radar, and digital photogrammetry);
- access to free open-source remote sensing and geographic data (with the ability to use free, open-source software);
- economic cost-effectiveness of remote sensing data in businesses (e.g., forestry);
- increased computing power; and
- increased knowledge about remote sensing and GIS by the public with access to sites such as Google Earth, and personal use of GPS.
The geospatial data era is well underway – for researchers, for governmental agencies, for businesses, and the private citizen – at the time of writing this compendium.

**Framework**

There are basic components involved in acquisition, processing and product assessment of remotely sensed data. If we simplify this in a diagram, it could be represented as shown below. This compendium will refer back to this diagram, when discussing the different elements and processes.

![Diagram of framework](image)

**Study questions**

1. What is a good definition of remote sensing? Can you add to the definitions given in this chapter?

2. Do you think the definition of remote sensing will change over time?

3. In which situations might it be good to use remote sensing?
2. PHYSICAL BASIS OF REMOTE SENSING

Electromagnetic spectrum. Electromagnetic radiation is magnetic and electric fields that travels in the form of waves, with the speed of light. The spectrum stretches from gamma rays with $10^{-11}$ m wavelengths over visible light with 400 – 700 nm waves, to radio waves that are more than 1 m long.

Irradiance (E). Radiant flux from all directions, per unit area (W/m²).

Radiance (L). Radiant flux per unit area of surface per unit solid angle (W/m² Sr).

Reflectance ($\rho$). The ratio between incoming and emitted flux, observe however that the flux might be emitted differently in different directions, of which only one, or a few, directions are measured as radiance registered by the sensor. Reflectance is always between 0 – 1.

Reflectance factor (R). The ratio between the actually measured radiance from a target at a given direction and illumination situation and the radiance that is measured at a reference target that reflects fully and equally in all direction. The reflectance factor is related to reflectance, but easier to measure, and might vary between 0 – $\infty$.

The definition of remote sensing can be quite broad, as it is simply based on the act of observing objects from far away. In theory, one could “remotely sense” objects based on several different techniques, for example sound waves (e.g., sonar). In this course, we concentrate primarily on remote sensing which makes use of recording the properties of electromagnetic radiation being reflected from objects on the Earth. In remote sensing, there are different qualities of the electromagnetic radiation recorded by sensors that are of importance, such as “how much?” and “when?” An example of “how much” can be the intensity level of spectral data, while an example of “when” may refer to the timing of a laser pulse. To understand how remote sensing works, it is helpful to have some definitions and to understand some basic laws of physics. This chapter introduces the basic properties of light and other electromagnetic radiation, and in particular how sunlight interacts with the atmosphere and the target. Thus, the emphasis is on factors that influences the spectral properties of “2D” images, like satellite images and air photos. How 3D data are created from photogrammetry or laser scanning, will be explained in later chapters.
2.1. The electromagnetic spectrum

Electromagnetic radiation can be understood both with wave theory and with quantum theory. In quantum theory, the energy of a photon or a quantum is given in Equation 2.1:

\[ Q = h \cdot \nu \]  

(2.1)

where

- \( Q \) = the energy of a quantum in Joules (J)
- \( h \) = Planck’s constant \( 6.626 \times 10^{-34} \) J sec
- \( \nu \) = frequency of the energy

One obvious source of electromagnetic radiation is the sun, but there are many other sources as well. All objects with a temperature above absolute zero emit energy, and the energy emitted is in proportion to its temperature, which will be described later.

Electromagnetic radiation has both electric and magnetic field components (seen in Figure 1.2 in Lillesand et al.). The radiation travels through a vacuum at the velocity of light, \( c \) (where \( c \) is a constant equal to \( 299,792,458 \) m/s or for simplicity \( 3 \times 10^8 \) m/sec).

The time point at which there was a breakthrough in understanding light’s electromagnetic properties came in 1864, when James Clerk Maxwell published his paper “A dynamical theory of the electromagnetic field”. Maxwell derived a wave form of the electric and magnetic equations, thus uncovering the wave-like nature of electric and magnetic fields, and their symmetry. Because the speed of the electromagnetic waves predicted by the wave equation coincided with the previously determined and measured speed of light, Maxwell concluded that light itself was an electromagnetic wave. It followed thereafter that

\[ c = \nu \cdot \lambda \]  

(2.2)

where \( c \) = speed of light, \( \nu \) is the energy wave frequency, and \( \lambda \) is the energy wavelength. Wavelength is the distance between successive wave peaks and frequency is the number of cycles passing a fixed point in a given period of time (see Fig 1.2 in Lillesand et al.).

By combining equations (2.1) and (2.2) we get:

\[ Q = hc / \lambda \]  

(2.3)

Therefore, what is useful to remember is that the energy of electromagnetic radiation is inversely proportional to the frequency of the radiation, or in other words, short wave radiation, for example blue light, will generally carry more energy than longer waves.
Note that since the wavelength and the frequency are inversely proportional to each other, only one of them needs to be stated to describe the wave. In some cases, frequency is used to describe the wave (as in radar remote sensing) and in other applications (as in optical remote sensing), the wavelength is most often used. For convenience, the wavelengths can be ordered and arranged along the electromagnetric spectrum (Figures 2.1, 2.2). Significant ranges of wavelengths have been categorized by giving a name to that range, for example, the ultraviolet spectrum, visible spectrum, infrared spectrum, microwaves, and radio, to name a few. Some spectral ranges of importance are named Table 2.1.

Table 2.1 Examples of electromagnetic wavelength bands used in remote sensing*

<table>
<thead>
<tr>
<th>Name</th>
<th>Approximate</th>
<th>Actual</th>
<th>Use in remote sensing</th>
</tr>
</thead>
<tbody>
<tr>
<td>Visible spectrum</td>
<td>0.4 – 0.7 μm</td>
<td>0.380 - 0.750 μm</td>
<td>Some optical sensors</td>
</tr>
<tr>
<td>- Blue</td>
<td>0.4 – 0.5 μm</td>
<td>0.450 – 0.495 μm</td>
<td>Most optical sensors + LiDAR for water depth</td>
</tr>
<tr>
<td>- Green</td>
<td>0.5 – 0.6 μm</td>
<td>0.495 – 0.570 μm</td>
<td>Most optical sensors</td>
</tr>
<tr>
<td>- Red</td>
<td>0.6 – 0.7 μm</td>
<td>0.620 – 0.750 μm</td>
<td>Most optical sensors</td>
</tr>
<tr>
<td>Near Infrared light</td>
<td>0.7 – 1.4 μm</td>
<td>0.78 – 1.3 μm</td>
<td>Some optical sensors + many topo LiDARS</td>
</tr>
<tr>
<td>Shortwave Infrared light†</td>
<td>1.3 – 3.0 μm</td>
<td>1.3 – 3.0 μm</td>
<td>Some optical sensors + some topo LiDARs</td>
</tr>
<tr>
<td>Thermal Infrared</td>
<td>3.0 – 14.0 μm</td>
<td>3.0 – 14.0 μm</td>
<td>Sensed as heat, sometimes used in a separate sensor</td>
</tr>
<tr>
<td>Microwave</td>
<td>1 mm – 1 m</td>
<td>1 mm – 1 m</td>
<td>Used in radar</td>
</tr>
</tbody>
</table>

* It is useful to know that there are differing opinions on exactly where to draw boundaries and names for different ranges of wavelengths. See for example the “International Commission on Illumination” or different ISO standards or different textbooks.

† Note that some texts use 1.4 μm as an upper limit for NIR/lower limit for SWIR.

† Note that “Shortwave Infrared” is sometimes also referred to as “Mid-Infrared”, depending on the field of study.

There are other categories (listed below) that are also of significance in remote sensing. Note that these are not necessarily mutually exclusive, and that the categories are often defined for convenience and relevance to certain applications. Also note that there is sometimes confusion about use of the name of these categories, and you may see mistakes on the internet and other places.

- **optical spectrum** (0.30 – 15.0 μm) are the wavelengths that can be reflected and refracted with lenses and mirrors.

- **reflective spectrum** (0.38 – 3.0 μm) are the wavelengths whose reflectance can be measured, and are commonly used in “optical” remote sensing.
- **emissive spectrum or thermal spectrum** (3.0 – 100.0 µm), which are wavelengths which emit heat, and are measured by thermal remote sensing instruments.

The Electromagnetic spectrum is shown in Figure 2.1. Table 2.1 gives wavelength measurements in micrometers (µm), while Figure 2.1 gives the wavelengths in nanometers (nm). One µm is 1000 nm. We said that all objects emit some amount of radiation. How much? Why? The amount of energy an object emits is partly a function of the surface temperature, which is expressed by the Stefan-Boltzmann law which states that blackbody radiators (which is a hypothetical “ideal” radiator - an object which absorbs all and re-emits all of the energy incident upon it) have an amount of emitted radiation or radiant exitance \( M_e \) which can be calculated as:

\[
M_e = \sigma T^4
\]  

where

- \( M_e \) = total radiant exitance (i.e., “emitted radiation”) from the surface in Watts (W)/m\(^2\)
- \( \sigma \) = the Stefan-Boltzmann constant (5.6697 x 10\(^{-8}\) W m\(^{-2}\)K\(^{-4}\))
- \( T \) = the temperature of the object’s surface in degrees Kelvin.

Since any material that has a temperature above absolute 0 will emit energy, the Stefan-Boltzman Law shows that it will emit energy in proportion to its temperature raised to a power of 4. We will first perceive this electromagnetic energy as heat, and then if the temperature rises, also as light. For example, a piece of iron that is being heated will be red at a certain temperature and then become even white and maybe bluish. Why blue? There is a law for this phenomenon called Wien’s displacement law. It says that the dominating wavelength for the energy emitted from an object will be shorter and shorter as the temperature of the object rises, so that:

\[
\lambda_m = A / T
\]

where

- \( \lambda_m \) = the wavelength of maximum spectral radiant exitance in µm
- \( A = 2897.8 \) (or rounded up to 2898) µm K (a constant)
- \( T \) = temperature in degrees K

From Figure 1.4 in Lillesand et al., it is seen that 6000 K, which is the temperature of the surface of the sun, will result in a radiation maximum in the wavelength region of 0.4 – 0.7 µm, which is exactly the wavelengths the human eye can register! We also see that the normal temperatures of the Earth (around 300° K) will result in a radiation maximum in the wavelength region around 10 µm and this is in fact wavelengths that we perceive as thermal heat.
2.2. Interaction of energy with the atmosphere and Earth

To understand just what the remote sensor is recording, it is necessary to understand the interactions between light, the atmosphere, and the objects being observed. Figure 2.2 below illustrates the topics to be discussed in the following sections.
Figure 2.2 – Explanation of total radiance that is recorded by the sensor. The total incident radiation upon an object \( (E_{\text{tot}}) \) comes from a combination of the direct but attenuated sunlight \( (E_S) \) (“attenuated” means “reduced intensity”) and skylight (or diffuse irradiance, \( E_d \)) which hits an object and reflects with reflectance \( \rho \). The attenuated radiance reflected from the object (which is equal to \( \rho \cdot E_{\text{tot}} \cdot T / \pi \), where \( T \) = the transmission of the atmosphere) is combined with path radiance \( (L_p) \), to equal the total radiance \( (L_{\text{tot}}) \) recorded by the sensor (in other words, \( L_{\text{tot}} = (\rho \cdot E_{\text{tot}} \cdot T / \pi) + L_p \)).

2.2.1. Atmosphere

Let’s first understand the influence of the atmosphere on the radiation coming from the sun, as well as its influence on radiation reflected back from the objects on Earth. During optical remote sensing of the Earth’s surface, the atmosphere will interfere with the light in several different ways.

One important mechanism is that incoming and outgoing electromagnetic radiation (light) is absorbed by molecules in the atmosphere (such as carbon dioxide, oxygen, ozone). In this process, shortwave light energy is converted to longer-wave heat energy. Different gas molecules interact with different wavelengths of the electromagnetic spectrum. For this reason, there are only certain “windows” (or wavelength bands) that are open for remote sensing through the atmosphere (Figure 2.3).
Consequently, remote sensing sensors are most often designed to operate within these “atmospheric windows” where absorption of light energy is minimal.

![Atmospheric transmission in different wavelengths](image)

Figure 2.3 Atmospheric transmission in different wavelengths shown in purple. The transmission is zero in several wavelength areas due to gas absorption. Figure from Wikipedia.

The other major mechanism present in the atmosphere is scattering, which means that light interacts with molecules in the air and changes direction. The way in which light scatters is related to the wavelength, but in a more continuous way than with absorption. Scattering is also most apparent for shorter wavelengths, such as ultraviolet and blue light, and to some degree in the green and red wavelengths.

There are three main scattering mechanisms:

- Rayleigh scattering
- Mie scattering
- Aerosol scattering

**Rayleigh scattering** is caused by small molecules in the atmosphere. Shorter wavelengths (blue and violet) are scattered much more, because the strength of the Rayleigh scattering mechanism is roughly proportional to the wavelength. This is the main mechanism that causes the sky to look blueish. Since Rayleigh scattering is due to interaction with air molecules, barometric pressure can be used to estimate the amount of Rayleigh scattering.

**Mie scattering** is caused by atmospheric particles of about the same size as the wavelength of the light.

**Aerosol scattering** (or nonselective scattering) is caused by larger atmospheric particles (dust, haze, smoke, etc). Aerosol scattering has a larger day-to-day variation than the other scattering mechanisms. Since it is
caused by large particles near the ground, aerosol scattering will be dependent on the weather situation, as well as the amount of air pollution in the area.

Scattering and absorption of energy can occur on the path from the sun to the Earth. When it is scattered and then hits the target it is referred to as sky irradiance or diffuse irradiance. Refer back to Figure 2.2 and observe that the main radiation components reaching the ground are a combination of direct sunlight and diffuse irradiance. The sum of these two components upon an object is called incident radiation.

2.2.2. Energy interaction with objects on Earth

What happens to the incident energy that hits an object? Total incident energy = reflected energy + absorbed energy + transmitted energy. In other words, no incoming energy will “disappear” as it interacts with an object; all of it is either reflected, absorbed or transmitted. An illustration of this is given in Figure 2.4. The reflected light is what is seen by our eyes and recorded by sensors (e.g., cameras, satellite sensors, etc.).

![Energy interaction diagram](image)

Figure 2.4 The incident energy which meets the leaf is either absorbed and used in photosynthesis, or is transmitted or reflected. (adapted from Figure 10:3 in F&F).

2.3. Reflectance and Reflectance factor

“Reflection is when incident energy (incoming light) hits an object and bounces off. The color of an object is actually the wavelengths of the light reflected while all other wavelengths are absorbed or transmitted. The physical and chemical composition of matter determines which wavelength (or color) is reflected.” - NASA

Reflectance (denoted as “p”) is easy to define in theory:
Reflectance = reflected energy / incident energy
Reflectance (ρ) is always given as a number between 0 and 1. However, to measure reflectance is more difficult, since both the incident radiation (coming in from all directions), and the outgoing radiation (going out in all directions), need to be figured into the calculation. In practice, a concept known as the Reflectance factor (R) is therefore used more often than Reflectance (ρ).

The Reflectance factor (R) = the radiation from a target that is actually measured from a given position for given illumination conditions, divided by the radiation that would have been measured if the target had been an ideally reflecting Lambertian surface (described in next section). In practice, the Reflectance factor can be measured by making a radiometer measurement over specific objects of interest (for example, a small plot of crops, or grass). Then a plate made of a material with near Lambertian properties is placed in front of the radiometer instead and a second radiometer measurement is made. The Reflectance factor is the ratio between the first and the second radiometer reading. Different objects have different reflectance factors, depending on the physical and chemical composition of the matter (as the NASA statement above says).

The proportions of reflected light in different wavelengths causes color. It is however only the diffuse part of the reflection, also called Lambertian reflection (= that reflects equally in all directions), that carries color. The opposite of Lambertian reflection is specular reflection. This means that the light is reflected like a mirror, (the angle of incidence has an equal size as the angle of reflection). Most objects in nature, such as vegetation, have properties that are a mix between ideal specular reflectors and ideal Lambertian reflectors. Figure 2.5 shows an example of Lambertian reflection and specular reflection.

| Reflectance factor (R) = the radiation from a target that is actually measured from a given position for given illumination conditions, divided by the radiation that would have been measured if the target had been an ideally reflecting Lambertian surface (described in next section). In practice, the Reflectance factor can be measured by making a radiometer measurement over specific objects of interest (for example, a small plot of crops, or grass). Then a plate made of a material with near Lambertian properties is placed in front of the radiometer instead and a second radiometer measurement is made. The Reflectance factor is the ratio between the first and the second radiometer reading. Different objects have different reflectance factors, depending on the physical and chemical composition of the matter (as the NASA statement above says). The proportions of reflected light in different wavelengths causes color. It is however only the diffuse part of the reflection, also called Lambertian reflection (= that reflects equally in all directions), that carries color. The opposite of Lambertian reflection is specular reflection. This means that the light is reflected like a mirror, (the angle of incidence has an equal size as the angle of reflection). Most objects in nature, such as vegetation, have properties that are a mix between ideal specular reflectors and ideal Lambertian reflectors. Figure 2.5 shows an example of Lambertian reflection and specular reflection. Figure 2.5. Demonstration of Lambertian reflectance (left) and Specular reflectance (right). 2.3.1. Calculation of radiance received by a sensor It is only a few percent of the light energy hitting the Earth that is reflected back up to the atmosphere. The reflected light that reaches the
sensor is also combined with the path radiance, which is radiation that comes from scattering of light in the atmosphere, and that have reached the sensor without having been reflected from the target.

Formula (Eq 2.6) show how to calculate what is being measured by the sensor. The solar radiation that reaches the Earth is measured as the total incident radiation per area unit, and called Irradiance, which in physics is abbreviated (E) and has the unit measure Wm\(^{-2}\). The two main components of Irradiance (E) are the diffuse irradiance from the sky (Ed) and the direct sunlight (Es). These two components are often summarized to a measure of total irradiance (E\(_{\text{tot}}\)) hitting the target: (E\(_{\text{tot}}\) = Ed + Es).

A sensor measures the radiation that is received from a given direction, in a cone angle. The sensor measures Radiance, which in physics is abbreviated (L) with units Wm\(^{-2}\) Sr\(^{-1}\) where Sr stands for steradian, which is the unit of a cone angle.

The total radiance reaching a sensor can be written:

\[
L_{\text{tot}} = (\rho * E_{\text{tot}} * T / \pi) + L_p
\]  
\[ \text{L}_{\text{tot}} = \text{Total radiance (Wm}^{-2}/\text{Sr}) \text{ reaching the sensor} \]
\[ \rho = \text{Reflectance for the studied object} \]
\[ E_{\text{tot}} = \text{Total irradiance hitting the studied object at ground} \]
\[ T = \text{Transmission in the atmosphere (0 – 1)} \]
\[ L_p = \text{Path radiance (Wm}^{-2}/\text{Sr}) \text{ which is a radiance component from the atmosphere that reaches the sensor without having reached the target.} \]
\[ \pi = 3.14159… \text{ (the need for a } \pi \text{ in the formula can be derived from the fact that } E_{\text{tot}} \text{ considers Irradiance from all directions and } L_{\text{tot}} \text{ measures radiance from one direction).} \]

The formula in Eq 2.6 could be written for a defined wavelength region (e.g., for example visible light from 0.4 – 0.7 µm). However, it is most usual to use a general “radiance per unit wavelength” term instead. This term is spectral radiance (L\(_{\text{λ}}\)) and the unit is Wm\(^{-2}\) Sr µm.

### 2.4. Practical implications of atmospheric influences

Sky irradiance leads to the shorter wavelengths of light, particularly the blue light, coming from all directions in the sky and not just from one direct path straight from the sun. This has important implications for aerial photography as well as for satellite remote sensing. One such implication is positive, since sky irradiance makes it easier to obtain information from the shadowed parts of an image, e.g. “between the trees”, if the blue light is present. Thus a “true colour” (an image with blue, green and red wavelengths – the wavelengths our eyes see) image is better for this purpose than a “false colour” image with a near infrared band but no blue
A further implication is that when taking photos from high altitudes, there will be a lot of blue light reaching the sensor which is reflected from air molecules and not necessarily from the object or ground. This will cause high altitude true colour photos to have a bluish haze, which is a negative effect.

The effects of the atmosphere can be seen in satellite imagery when one looks at viewing radiometrically uncorrected remote sensing images (i.e., without an atmospheric correction) or by inspecting the image histograms (image histograms will be discussed later in the compendium. We haven’t covered the subject of histograms yet, but for now remember that histograms are simply a plot of the frequency of the values of a chosen variable). The effect of a thick atmosphere compacts the histograms primarily in the short wavelengths (e.g., blue wavelength the most), resulting in a very poor dynamic range. Path radiance adds a background brightness which shifts the histograms to a higher value range, again with a stronger effect in the short wavelengths. True-colour composites will reveal a bluish haze over the image with very poor contrast. Under clear atmospheric conditions, shadows will be much deeper, water will appear darker and there will be much better overall contrast. The short visible wavelengths, such as blue and green will be most severely affected by poor atmospheric conditions while the shortwave-infrared bands are least affected and typically have dark shadows and good contrast even in relatively poor atmospheric conditions.

The main problem with optical remote sensing, both using satellites and aircrafts as platforms, is that clouds make most days useless for image acquisition. This is one of the main advantages for radar sensors which operate with cloud-penetrating wavelengths. However, even if images are acquired during an apparently cloud free day, it should be remembered that the optical thickness of the atmosphere varies from day to day due to different weather conditions.

There are computer programs that for given atmospheric conditions can estimate $E_s$, $E_d$, $T$ and $L_p$. The most well-known of these programs is “6S” and also “ATCOR”. Theoretically, we could then solve the relationship between radiance registered by the sensor ($L_{tot}$) and the reflectance properties of the studied surface, $\rho$. However, as we will see later in the course, there are many uncertainties in such a procedure, and there are other, simpler methods to take care of atmospheric influences which are often “good enough.” However, this is an advanced topic in remote sensing.

3. Reflectance from boreal forest

Everything in nature has its own unique distribution of reflected, emitted, and absorbed radiation. These spectral characteristics can—if ingeniously exploited—be used to distinguish one thing from another or to obtain information about shape, size, and other physical and chemical properties. (Parker and Wolff, 1965, p. 21)
The sensor response in different wavelength bands will produce colour images that are used both for visual interpretation and automated classification and estimation of landcover classes or forest variables.

Living vegetation will absorb blue and red light, but absorbs less green light, which is the reason why vegetation appears green for our eyes. If our eyes would have been sensitive also for near infrared light (= the light just beyond 0.7 µm) we would see that the reflectance from living vegetation is particularly high in this wavelength region (Figure 2.6). This phenomena is used in color infrared images, where the sensor response from the near infrared light is displayed as red. The typical reflectance curves shown in Figure 2.6 will however be less distinct when the observed area change from radiometer measurements of leaves, to image pixels of branches, full trees, or groups of the trees including their shadows. The color of a 10 x 10 m satellite image pixel in a forest will be composed of four components: sunlit canopy, sunlit ground, shadowed canopy, and shadowed ground. The fraction of shadows is an important factor for the brightness of a forest pixel, in particular at our northern latitudes, where the sun angles are low. In fact, there is a correlation between how dark a forest pixel is and the stem volume (or correlated variables like tree biomass or basal area). This might be explained by the fact that large trees, or many trees, causes more shadows than small or few trees. This correlation between stem volume and low pixel values, is stronger with longer wavelengths. The reason for this is probably that the shadows are darker in the longer wavelength bands where there are less scattered light from the atmosphere. The exception to this general rule with gradually improved correlation with stem volume with longer wavelength bands is the near infrared region. In this region the correlation with the photosynthetic activity within the pixel will be stronger than the correlation with the tree biomass.

![Figure 2.6. Typical reflectances from vegetation, soil and water, as a function of wavelength.](image-url)
Study Questions

1. Blue light is scattered more in the atmosphere than longer wavelengths such as for example red light. Which advantage and which drawbacks does the scattering of blue light cause when using aerial photography?

2. In boreal forests, areas with high stem volume will appear as dark in most wavelength bands when analyzing optical satellite images. Mention a possible explanation for this.
3. FOUNDATIONS IN REMOTE SENSING

Optical sensor: The term is used here to collectively define sensors which register the reflected sunlight, for example, aerial photography, airborne optical scanners, and satellite-borne electro-optical sensors.

LiDAR: A system which measures distances by registering the time between sending away a laser pulse, and when it is returned. A scanning system is often used, and therefore it is often referred to as laser scanning.

Radar: A system which creates an image through registration of how a returned microwave wavelength is returned.

Passive sensor: A sensor which registers reflectance whose source radiates from nature (e.g., reflected sunlight, or warmth radiated from an object on earth), as opposed to a sensor which sends out its own signal. Examples are aerial photography cameras, common multi-spectral optical sensors placed on satellites, and thermal cameras.

Active sensor: A sensor which sends out its own signal which is then also recorded by the sensor. Examples are LiDAR and radar.

This chapter is an overview and introduction to sensors commonly used to perform remote sensing of forests, which platforms are commonly used, and what type of data are produced by these sensors. A short introduction to forest information collected by these sensors is also given, but more details will be given in the individual chapters about the different remote sensing data types.

3.1. Sensors

Remote sensing can be performed using a wide range of sensors. We can categorize these sensors by their properties, often defined by the mechanism they use or the wavelength in which they operate. The following three categories – Optical, LiDAR, and Radar – are those currently most used in forest remote sensing today.

3.1.1. Active and passive remote sensing techniques

One important distinction between sensors is identification of them as passive or active remote sensing techniques. Passive techniques mean that the sensor registers energy that is either reflected by or emitted from an object. Active sensors send out energy themselves and record the signal that comes back. One example of an active sensor is the laser sensor which sends out very controlled light pulses and record the time delay and intensity for the reflected light pulse. Another important example is
radar, which sends out very controlled radio signals and creates an image from the returned radio signal. Table 3.1 exemplifies some passive and active remote sensing techniques and comments on their usability for forestry.

Table 3.1 Examples of passive and active remote sensing techniques and comments on their usability for forestry.

<table>
<thead>
<tr>
<th>Passive techniques</th>
<th>Optical wavelength region</th>
<th>Microwave wavelength region</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aerial photography and optical satellite sensors – Traditionally the most important remote sensing techniques for forestry. Today has the photographic film in aerial cameras been replaced with electro optical sensors such as CCD arrays and matrices working in the visible and near infrared wavelength regions. Using satellites as platforms, similar sensors are of great importance for also obtaining overviews of forest resources over large areas, and for detection of changes. A pixel value provide a measure of reflected light. Digital surface models can also be computed from overlapping images taken from different observation points.</td>
<td>Passive microwave – There are sensors on board satellites and aircrafts that registers the radio waves emitted from the earth, however due to the low energy emitted, the footprint (pixels size) is so large that these sensors have no practical importance for forestry.</td>
<td></td>
</tr>
<tr>
<td>Laser scanning – Laser systems work with light, often near infrared light. Distance measuring laser scanners are used for measuring digital elevation models, as well as for measuring tree heights and canopy structure. The produce a 3D point cloud which can be used for estimation of forest variables.</td>
<td>Radar sensors – There are many radar systems, both onboard aircrafts and satellites, depending mainly on wavelength some are less relevant for forestry, whereas other systems have a quite good potential. Especially systems with very long or very short wavelengths have a potential for forestry. The pixel value basically shows the electrical properties of the site of the pixel, but registrations from different view point can also be combined to form a digital surface model.</td>
<td></td>
</tr>
</tbody>
</table>

3.1.2. Optical sensors

In passive sensors which measure reflected electromagnetic energy, light is directed through a lens system to sensors that convert the light to electric pulses. There are different mechanisms for scanning the earth from an aircraft or satellite. The oldest principle is to have a rotating mirror in front of the sensor that direct swaths of light across the scene into the sensor. A
more modern technique with less mechanical parts is to use an array of sensor elements, often of a type that is called a CCD array (the inventors of the CCD array received the Nobel prize in Physics 2009). The sensor array senses a line of pixels at a time, and is dependent on the forward speed of the platform in order to read data from the next line which might be only 0.03 seconds later. A third version is to replace the film in frame camera with a matrix of sensor elements which instantaneously registers the earth.

The sensors usually record energy from specific bands in the visible and near infrared wavelengths. Some satellite sensors like for example Landsat 8 OLI also record reflected light in the mid infrared spectrum. The design is often based on what the intent of data collection is (e.g., for ocean, or land, or ice), but also what is technically possible, avoidance of certain wavelengths affected by the atmosphere, as well as cost. The sensors may record energy in a few wavelengths (e.g., the four bands blue, green, red, near infrared is a common choice). When a sensor records energy in many narrow ranges (e.g., 260 bands) it is called a hyperspectral sensor.

An example of an optical sensor can be found as close as your own personal camera. This records reflected light in the visible range. Another type of camera is the large-format camera, often used for aerial photography. These often extend into the near-infrared range of the spectrum. More details on optical sensors are given in Chapter 4 on Optical Satellite data and in Chapter 5 on Aerial Photography.

3.1.3. LiDAR

Laser scanning has been established as a new and effective approach to forest data capture. Airborne laser scanning measures with decimeter accuracy the location of points on the ground and in the canopy. Airborne laser scanning uses the laser's ability to measure distance using the time from when a laser pulse is sent out, until the laser light is reflected from the ground or vegetation comes back to the sensor. The technology to measure distances with laser is often called LiDAR from the English “Light Detection And Ranging”. More details on LiDAR are given in the Chapter 6 on LiDAR.

3.1.4. Radar

Radar (acronym for RAdio Detection And Ranging) is an active sensor for detecting, locating, tracking, and identifying objects even at a considerable distance. Radar is a technique that is not influenced by cloud coverage. More details about Radar are given in the Chapter 7 on Radar.
3.2. Platforms

The sensors used in remote sensing often need to be placed on a larger carrier which can operate to give a good position over the objects of interest. The carrier of the sensor is referred to as a “platform” and is a separate component of a remote sensing system. It may be anything from a satellite, to an airplane, to a stative. There are categories referring to the position of the platforms, namely spaceborne, airborne, and terrestrial. The choice of platform is based on the purpose of the sensor. For example, if larger areas need to be imaged, then the sensor needs to be high above the landscape, requiring a spaceborne or high-flying airborne platform.

3.2.1. Satellites

Satellite platforms are a common carrier of sensors, providing stable and repetitive measurements of the Earth. There are satellites used for communication, GPS, and many other purposes. But here we concentrate on what are called “Earth Observation (EO) satellites”.

The orbit characteristic refers to the movement or placement of the satellite in relation to the Earth. There are two main types of orbits used for remote sensing satellites:

- Polar (or “near-polar”) orbit – The satellite travels north to south over the poles and images the area of the Earth under it, which varies due to the rotation of the Earth. The altitude is typically between 600 km and 2000 km above the earth. When the satellite crosses the equator at the same time with every orbit, this is called a “sun-synchronous polar orbit”, which is very common for most EO satellites.

- Geo-stationary orbit – The satellite travels over the equator at a distance of 36 000 km at the same speed and direction as the rotation of the Earth, allowing it to always image the same area. There is a system of weather satellites that take images from the geostationary orbit, but the images produced from that great distance has not good enough resolution for forestry.

The electricity needed for operating the sensor and sending the data to earth is obtained from solar panels. Most satellites also have small rockets that are used for correcting the orbit when needed. The fuel for these rockets is often a factor that limits the lifetime of a satellite. A typical lifetime is around five years, but some satellites have been operated much longer.

3.2.2. Airplanes/Helicopters

Aerial photography is often made with purpose-specific planes that have one or two central openings in the floor of the plane. These openings have
standardized dimensions where the aerial cameras or other instruments, e.g., a laser scanner, can be placed. A "normal" flying altitude for aerial photography is usually between 2000 m and 4500 m. Even the acquisition of “high altitude images” may occur at around 10 000 m flying altitude.

Aerial planes can also be used for low altitude photography around 1000 m flying altitude, but for this kind of shooting, often done for smaller rather than larger area coverage, there are also several other options. Photography or scanning at low altitude can be made by several companies that have cameras, or laser scanners, mounted on the outside of sport airplanes or helicopters.

### 3.2.3. Drones or UAVs

UAV is an abbreviation for Unmanned Aerial Vehicle and a common synonym is drones. Other common names are model aircraft, UAS (Unmanned Aerial Systems), or RPAS (Remotely Piloted Aircraft Systems).

Since the early 2000s UAVs have been used for forest applications, but it is only in recent years (2010s) that it has had an impact. A contributing factor to the breakthrough is that the systems have become cheaper, easier to fly and there is small cameras and software that makes it easy for the user to create orthophotos and 3D point clouds from images.

### Regulations

In Sweden, the Swedish Transport Agency issues flight rules regarding drones. The main limitation of the current regulations for forestry purposes is that the flight must take place within the operator’s sight. The drone platforms are divided into categories based on the starting weight and impact energy. A heavier platform requires higher security and is limited to a maximum flying altitude of 120 meters above ground level.

Permission needs to be sought from the Transport Agency for flying on missions, research or testing, but not if you are flying as a private individual for non-commercial purposes. The regulations must be followed even if you do not have to seek permission. Read on the Transport Agency website (www.transportstyrelsen.se) for the current rules. Besides permits for flight, you may have to seek permission for taking and dissemination of any photos acquired from a UAV. However, the rules around this are currently changing in Sweden and it is recommended to check up to date information.

### Safety

There are often a number of safety features built into the autopilot system that can be triggered by various events that can, in turn, compromise security. Examples of such triggering events are

- Flying off the defined flight range (geo-fence)
- Loss of radio contact (returning and landing)
- Loss of data link
The action to be taken when a triggering event occurs, the pilot can usually take some counter-action. Here are some examples of actions:

- Return to the starting point and land
- Continue the mission despite the incident
- Reconnect the manual pilot function and allow the pilot to take over the flight.

### 3.2.3.1. UAV Airplane

An airplane or flying wing has a technically simple design. Figure 3.1 shows a flying wing UAV from the Swedish manufacturer SmartPlanes. The sensors are usually mounted inside the fuselage, which also provides protection on landing.

![Figure 3.1](image-url)

Figure 3.1 This is a typical setup when flying. The pilot performs a pre-flight check. Data Modem (1) communicates with the aircraft (2) in real time and via the ground station software in the computer (3) the pilot can obtain information on the plane's location, speed, incline, etc. The remote control (4) is used to control the plane during takeoff and landing, but at any time during the flight the autopilot can be used to manually control the aircraft.

**Control**

Control can be done completely manually through a standard remote control, but usually tend to make use of the autopilot stabilization at takeoff and landing. With the help of the autopilot function, the plane can also take off and land automatically. How the start and take off are performed will vary between manufacturers. Some deliver a startup ramp, while others require the user to throw the plane into the air.
Advantages
There are a number of aircraft weighing less than 1.5 kg and thus fall under the Transport Agency’s “Class 1A”, which has no limit in flying altitude (however, must always be in the pilot’s sight). The fuselage is often manufactured in polystyrene and is thus a very cheap and durable construction. It can even land without the engine turned on because it can glide to the landing. The sensor is usually mounted inside the fuselage and is well protected by a hard landing or crash. The platform can be flown without the support of the autopilot.

One advantage of an airplane UAV platform is that you do not need any electronics (stabilization) to be able to fly, but it is enough that the pilot can control two servos. Even if these were to malfunction that does not necessarily mean that the aircraft would crash, and that the sensor would be destroyed.

Because the wings provide lift capability, very little battery is required. This gives a long flight time. Currently the regulations are more of a limiting factor rather than the battery time to fly larger areas.

Disadvantages
The speed depends largely on the wind and one should plan missions with the wind coming from the side of the flight direction in order to get the most constant possible air speed. However, one problem with the plane platform is the constant lean and therefore you will not be taking photos in nadir orientation if you do not have a gimbal mounted. During takeoff and landing, a relatively large area is required, for example, a clearing or a field. For the experienced pilot take-off and landing can occur on a road (untrafficked, of course!) even though the trees are close.

3.2.3.2. Multirotor
A multirotor platform is able to lift straight up by pushing air downwards. This technology has been further developed in recent years (2010s) thanks to cheaper inertial navigation systems. To keep a multirotor platform stable the motors must be controlled with high precision. If the vehicle starts to tilt on take-off, one or more motors must change speed to correct for this. There are a plethora of manufacturers and Figure 3.2 shows an example of a quadrocopter from the American company 3D Robotics.
Figure 3.2 Quadrocopter (1) of 3D Robotics model Solo. On this platform, there is a Sony Alpha SLR 5100 camera (2) with a 20mm lens.

Control
A multirotor UAV platform assumes that you have some kind of stabilization or autopilot that adjusts fan speed as the platform stays level. Many autopilots help the user to have control by simplifying the procedure, e.g., by keeping the platform stationary using inertial navigation (INS) and GNSS (GPS). Control can be done via both the autopilot and manually via the remote control.

Advantages
Take-off and landing goes very fast because the craft lifts straight up and can quickly get up to altitude where data collection occurs. The flight assignment (mapping) can be done quickly thanks to the fact that multirotor platforms do not need to fly a circle at the turn otherwise made at the end of each flight path. The speed can be kept constant, however, there will be a different pitch, but this can be solved with a gimbal.

Disadvantages
Battery time is in the current situation one of the major constraints and depending on altitude, you cannot cover such areas. Another disadvantage is that you basically cannot use a platform which will be in the Transport Agency's Class 1A; this is due to the weight being higher than 1.5 kg with the camera.

With a motor malfunction, the quadrocopter will generally crash, but a hex- or octocopter may be able to fly despite an engine malfunction. The body is also very sensitive to hard landings or crash landings. The sensor is usually
mounted hanging under the craft and is thus easily damaged during hard landing or crash. A multirotor is completely dependent on autopilot or stabilization electronics and cannot fly without it.

### 3.3. Ground-based and mobile systems

**Ground-based systems**

Both laser scanning as well as images acquired with digital cameras can be made from sensors located on a stationary tripod standing on the ground. In this manner, the data can provide tree stem position and shape, etc. To get the 3D data from digital photogrammetry the photos must be acquired from more than one position, after which the images are co-processed with a special software. Even in laser scanning, there may be reason to scan from more than one position, so that the number of hidden trees are minimized (e.g., trees that cannot be seen because another tree stem blocks them from view). This type of ground-based registration has great potential, but is not yet operational in any significant way.

**Mobile systems**

It is also possible to record data about tree stems and positions via a mobile (moving) sensor located on the ground. A laser scanner can be mounted to a backpack, or on an ATV or a forest machine. The type of laser scanner suitable for this, however, does not give as accurate measurements as larger systems that are fixed to a tripod. It is also possible to acquire 3D information about the tree stems from a moving video recording. This technique is called "Shape from motion" (SfM).

When mobile data are acquired, we need to know the position of the sensor at any moment, which can be difficult in the forest where GPS signals provide limited positional accuracy. As a complement to GPS, inertial navigation systems are used, as well as techniques that use the trees depicted as reference points for the sensor movement, a technique known as Simultaneous Location and Mapping (SLAM).

### 3.4. Remote sensing data models

Digital geographic data can be represented in several formats, the primary types being vector and raster. Vector data comprises point, line, and polygon format. Raster data have a defined number of rows and columns, with a set number cells (see figure 3.3). The grid cells have a defined equal size and may be square or rectangular. Raster data can consist of a single raster, or there may be several rasters layered in a single file. A raster may also be referred to as a grid or a matrix.

Another characteristic of remote sensing data is dimensionality. We currently talk about data as having dimensions, and being two-dimensional (2D), three-dimensional (3D), and even four-dimensional (4D). Two-dimensional means that the data have East-West and North-South geographic coordinates, while 3D means that the data have X, Y, and Z.
(Elevation), and 4D refers to the previous three dimension, plus the dimension of time.

3.4.1. The principal data structure of a digital raster image

Let’s consider the structure of a digital raster image. All imaging devices discussed in this section produce a digital image which could be viewed as a matrix of grid cell values, where each element in the matrix corresponds to the radiance that the sensors measure from the corresponding area on the earth, e.g. 0.5 m × 0.5 m for a digital aerial photo, or 10 m × 10 m for a satellite image (Figure 3.1). The matrix element can store a value, for example, between 0 and 255 (since 256 values can be stored in 8 bits in the computer). When several wavelengths are acquired, these are represented by having one “layer” for each colour. The conceptual data structure described here (a matrix or raster) is identical to those raster data structures used in geographical information systems (GIS).

![Illustration of one band in a digital image](image)

Figure 3.3 Illustration of one band in a digital image, the value in each cell represents a measurement (usually reflected light), for the corresponding position on the ground.

Optical satellite data are often delivered in 2D raster format, as a file that consists of several raster layers, where each raster represents a different wavelength. For example, optical satellite data from a sensor which records reflectance in the green, red, near-infrared and short-wave infrared bands will have four raster layers. The rasters are geographically co-registered to each other, and often (but not always!) have the same grid cell size.

The raster grid cells in the optical satellite data are called Pixels, which is short for Picture Elements. Rasters from remote sensing can be thought of as image files, and often have a data format that stems from this background. They may be TIFF, GEOTIFF, JPG, BIL, BSQ, IMG, JP2000 format, among others. These formats are common, and it is possible to export from one format to another.
Aerial photographs are also taken in raster format and consist of pixels. In addition 3D point data can also be created using raster data as the source. This process is based on photogrammetry which is covered in Chapter 5.

Point data consist of a single point, without area, in geographic space. The point can have 2D coordinates, but can also be represented in 3D or 4D. Currently, in many remote sensing applications, 3D remotely sensed data are creating a revolution in information, particularly for forestry where knowing the height of trees is of great importance. LiDAR data are delivered as point data, with 3D coordinates. The may be delivered in -.LAS or -.LAZ file formats.

3.5. About data resolution

When describing different characteristics of remotely sensed data, there are different types of “resolution” that are important. Each sensor has its own characteristics of importance. You often need to know the characteristics of a sensor to know if it will be useful for different applications. The characteristics of active sensors and passive sensors are different, and we’ll first concentrate on passive sensors.

Whether or not we can see our objects of interest on the Earth, from above, is one of the important questions we ask ourselves when considering which remote sensing system to use. Can we see the individual trees or are we limited to only being able to identify groups of trees in stands?

In this respect it is a kind of resolution of the instrument we are interested in. In remote sensing, we can often talk about different kinds of resolution when we describe a sensor. The resolution characteristics are

- Spatial
- Spectral
- Temporal
- Radiometric.

The word resolution is used in different ways in remote sensing, which might cause some confusion. In photographic systems, it is used for the capability of a film and camera system to resolve a test pattern of black and white lines. In digital systems, it is often used as a synonym for pixel size, as it would be measured on ground. To avoid confusion, it is better to call this measure pixel size, or more correctly, ground sampling distance (GSD). Although there are many concepts that are similar between photographic film-based cameras and digital electro-optical sensors, it is good to note that the concept of “resolution” is different between these two systems.

The pixel size, based on the pixel spacing (the GSD) in an image is determined by the platform (and scanning) speed and the sampling rate. The GSD tells the distance between pixel readings, which might be
different along track and across track. It is also common that digital images have been resampled to a new pixel size through some pre-processing steps.

The area that the sensor actually “sees” at one moment is called the *ground resolution cell* (GRC). This area could be both slightly larger, and smaller than the area given by the GSD. Figure 5.4 in Lillesand et al. illustrates this concept.

The size of the GRC depends on the characteristics of the sensor as well as flying characteristics (height, look angle, etc). The GRC is calculated as the segment of the ground that is measured, which is circular and has a radius of \( D \), where

\[
D = H' \times \beta
\]

where

- \( H' \) = flying height above the terrain
- \( \beta \) = the Instantaneous Field of View (IFOV) of a system, which is expressed as an angle expressed in radians.\(^1\)

The radiance measured from the GRC is then recorded by the sensor. Depending on the properties of the sensor, the actual shape of the GRC area can be a circle or a square/rectangle.

*Image extent* refers to the area on the ground that a single image covers. The pixel size often has a relationship with Image Extent, in that large pixel sizes often allow for larger image extents whereas small pixel sizes tend to be collected over smaller image areas. In some cases the image extent can be described as having a limited area in both width and height (e.g., 60 km x 60 km), or in the case of some scanners, only a width is referred to, often called the *swath width*, as the scanner is continually moving in one direction and is only limited by the swath that it covers. Here, the data are often extracted according to the user’s needs and according to how the delivering company wishes to sell the data.

A combination of the sensor properties and the flying height of the sensor (in the case of the satellite sensor, the *height of orbit*) will play a role in determining GRC and Image extent.

*Radiometric resolution* sometimes refers to the quantization that is used by the sensor to record the incoming radiance. It indicates the sensitivity of the instrument. The radiometric resolution is often expressed in the number of bits used in the quantization. While many previous sensors have had 8-bit radiometric resolution (with values from 0 – 255), the new trend is towards higher radiometric resolution, such as 10- and 12-bit. The advantages of this will be discussed later. However, even if sensor can store data in 256 different level, the actual meaningful radiometric resolution might be less,

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\(^1\) Please refer to Lillesand et al. Figure 5.2 to see a graphic demonstration of this concept.
due to for example technical limitations in the sensor system. If for example one wavelength band in spectral sensor is very narrow (e.g having a high spectral resolution, see below), it might happen that the signal-to-noise ratio (SNR) will become low, which in reality will limit the actual radiometric resolution.

The spectral resolution for a wavelength band refers to the width (in nm) of the wavelength band sensed by the sensor. This is not just a matter of indicating whether they are blue, green, red, or Near-IR, but the specific wavelengths that the instruments are developed to sense. Hyperspectral sensors, for example, sense hundreds of narrow ranges of wavelengths, and are therefore said to have a high spectral resolution.

There is something called temporal resolution also, which refers to the revisit time of a sensor. For example, some satellites pass over the same spot on Earth every 16 days, while others may pass over every 5 days. The satellite that passes over more often has a higher temporal resolution (although, in reality is clouds often the limiting factor).

An interconnected characteristic is the view angle of the sensor. Some satellite sensors are constantly looking straight down, and acquiring what are called “nadir-pointing images”, meaning that at the mid-point of the image, the sensor is looking straight down, without any angle. In contrast, some sensors can acquire data by pointing their sensor at an angle, and these are called “off-nadir images”. The advantage to a pointable sensor is the flexibility of taking images in several areas. If it is cloudy at nadir, and a non-cloudy area is viewable at +20° viewing, then that image can be acquired instead. The ability to point the sensor allows for a higher temporal resolution.

3.6. Display of raster images

To understand the displaying of optical data from remote sensing, it is necessary to discuss some basic concepts. The human eye sees color only in the “visible wavelengths”, that is, the red, green and blue wavelengths. Remember that most raster files will contain multiple spectral bands or layers of data within a single image file. Often, there is one or more wavelength represented that we do not usually see with the human eye.

The displays that we use, for example on computer screens, are made for our human eyes, which view in the visible spectrum, consisting of colors from violet to blue to green to yellow to red. Most image processing and GIS software that is used to look at image data uses a display with 3 “color guns” – Red, Green, and Blue. A way of looking at the image data is to put one band and display it with a single color gun. Say for example, we choose to display the red wavelength data using the red color gun display. Remember that each layer is made up of a raster of digital numbers, whose values will be (usually) 8-bit, 10-bit or 12-bit or whatever the radiometric resolution of the image was. The software will use the Digital Numbers (DNs) in the raster to display this amount of red for that layer, in a gradient.
If you were to look at the red wavelength, in just the red band, it would be a red image of varying red hues.

There are three “color guns” for displaying. Therefore, we can add another wavelength into another color gun – say the Green wavelength into the green color gun display. And then the Blue wavelength into the blue color gun display. This makes things easy, and it gives us a sense of “true color”, because this is what our eyes would actually see. We could just as easily have done any combination we wanted to, such as putting the Blue wavelength in the red color gun display, and so on.

3.6.1. **Trichromatic theory of color vision**

The human eye has three types of rod cells: short-, middle- and long-wavelength sensitive (blue, green and red). The trichromatic theory states that when all three types are stimulated equally, we see white light. Other colors are perceived when the three rod cell types are stimulated with different amounts of light:

\[ C = (b_R \times R) + (b_G \times G) + (b_B \times B) \]

where \( C \) is the perceived color, \( R, G, B \) are the three primaries (red, green, blue) and the \( b \)'s are the proportions of each. To use the R-G-B primaries on a computer screen or a projector is an *additive* principle.

Other processes are fundamentally *subtractive*, such as most inks on white paper, which use the subtractive primaries Cyan-Magenta-Yellow. These primaries are complementary colors of red, green, and blue. The complementary color for each primary color is obtained by mixing the other two primaries (e.g. cyan is the complementary color of red, and is produced by adding red and green).

*Color additive theory* says that when we take the colors Red, Green, and Blue, and add one to another, we get new colors (cyan, yellow, magenta), and if we add them all together, in equal amounts, we perceive white (Fig 3.4).

![Figure 3.4 – A demonstration of the concept of additive color theory](image-url)
A Color Space is a coordinate system for measuring and specifying colors. Some color spaces are designed to Euclidean distances between coordinates represent approximately perceptual differences between colors. The RGB color cube (Fig 3.5) used in image display on computer monitors is defined by the tristimulus coordinates of the phosphor response.

All colors that can be realized by the display are represented by (R, G, B) coordinates in the cube:

![RGB color cube](image)

Fig 3.5. The RGB color cube.

In general, the R-G-B color space:
- is not perceptually uniform;
- cannot display all colors;
- is very device dependent.

A Red-Green-Blue coordinate triple for each cell determines the intensity of each primary color on the display. A large number of colors can be created by mixing the intensities of R-G-B primaries \((256)^3 \approx 16.5\) Million unique colors!

### 3.6.2. “True color” and “False color” images

Most aerial cameras record colour in four spectral bands: blue, green, red and near infrared. Often there is also a panchromatic band with a broader spectral bandwidth and smaller pixel size recorded. During later postprocessing of the images, the panchromatic band might be combined with the multispectral bands in order to obtain a “pan-sharpened” image with both colour and high resolution. The four colours above are sufficient for producing both true colour and false colour imagery (Table 3.2 below).

<table>
<thead>
<tr>
<th>True Colour</th>
<th>False Colour</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blue</td>
<td>Blue</td>
</tr>
<tr>
<td>Green</td>
<td>Green</td>
</tr>
<tr>
<td>Red</td>
<td>Red</td>
</tr>
<tr>
<td>Infrared</td>
<td>Infrared</td>
</tr>
</tbody>
</table>

Table 3.2. Standard schemes for displaying a true colour and a false colour infrared image on a colour screen.

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2 See also figure 7.36 in Lillesand et al.
A “true colour” image might be displayed from Landsat 8 data, by displaying B2 (blue wavelength) with the blue colour gun, B3 (green wavelength) with the green colour gun and B4 (red wavelength) with the red colour gun. A “true colour” image is not possible with SPOT data, since there is no blue band.

A “false colour” image where vegetation will have reddish colours for vegetation, such as in a colour infrared aerial photo, can be displayed using the following bands

<table>
<thead>
<tr>
<th>True colour (vegetation appears green)</th>
<th>False colour (vegetation appears red)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sensed wavelength</td>
<td>Displayed as</td>
</tr>
<tr>
<td>Blue</td>
<td>Blue</td>
</tr>
<tr>
<td>Green</td>
<td>Green</td>
</tr>
<tr>
<td>Red</td>
<td>Red</td>
</tr>
<tr>
<td>NIR</td>
<td>-</td>
</tr>
</tbody>
</table>

Often, in remote sensing we talk about displaying bands in R,G,B (Red, Green, Blue), so we may say, for example, that we have “displayed TM band 5,4,3 in R,G,B” (as the false colour display above).

### 3.7. Forest information from remote sensing data

Different remote sensing data sources are more or less appropriate given the application. To determine certain forest characteristics, you will need to choose one data source over another, and therefore should know the possibilities and limitations of different data sources. What can and cannot be determined, given the available data?

#### 3.7.1. Optical data

When using data in the visible and near-infrared spectral region, which is what is most commonly available from optical sensors, the main spectral components in a forest stand are:

- sunlit canopy,
- shadowed canopy,
- sunlit ground and,
- shadowed ground.

For example, if using satellite data with visible and near infrared as well as shortwave infrared wavelengths, the data from the blue, green, red, and the two middle infrared bands are highly correlated with each other, and the darkness seen in the pixels in these bands over forest areas will be mostly a
function of the size of the trees and the stem number, since shadows are a dominant factor behind the reflectance.

The Near Infrared (NIR) band is more correlated with the structure of the leaves and deciduous forest and grass have higher reflectance in this band than coniferous forest, or the reflectance coming from the field layer with lingonberry or blueberry. A difference, or ratio, measure between the Near Infrared band and the red band, (as well as other combinations of near infrared and the sum of the other bands), is well correlated with the photosynthetic activity. The thermal band is influenced by factors that are difficult to control, like moisture, and is seldom used for forestry studies.

Altogether, spectral data from optical satellites over boreal forest do not contain much more than two dimensions of uncorrelated data (as stated above: the NIR and all other bands), but since many forest variables also are correlated and related to different tree size variables, estimates of many more than two forest variables are often presented in remote sensing studies.

The shortwave infrared (SWIR) bands have been shown to be especially useful for forestry, since that wavelength region (around 2 µm) is sensitive to the amount of shadows, and thus to the size of the trees. The SWIR bands are the ones that are best correlated with forest biomass and thus also with forest stem volume. However, regardless of the choice of spectral band, the images will not be much darker when the stem volume becomes more than 300 m³/ha. We say that the signal is “saturated” at this point, and doesn’t provide the detail of information. In a similar way, the general dependence between stand age and spectral signature becomes weak after the first thinning cutting.

The ability to separate and classify coniferous and deciduous forests is well established in the remote sensing literature, as deciduous leaves reflect more of the NIR wavelength than coniferous trees (Lillesand et al., 2008). Mixed deciduous/coniferous forests are more difficult to classify correctly due to the spatial arrangement and uneven mixtures of the deciduous and coniferous species within pixels. For boreal and managed coniferous forests, optical satellite data have a primarily negative correlation with wood volume in the visible and mid-infrared bands. Correlation with the near-infrared band is more varied and sometimes there is no correlation. Forest canopy self-shadow has a large effect on the spectral response, yet it can be used to help derive stand parameters. For Swedish forest conditions, shadow is particularly important due to several factors including that coniferous tree species’ crowns tend to cast more shadows than deciduous species and that the low sun angles occurring at Sweden’s relatively high latitude produce more shadow and illuminate ground vegetation less. The SWIR bands have been shown to be of significance in forest parameter estimation, most likely due to their sensitivity to shadow patterns.
As forests mature, the canopy tends to close which results in a weakening ability to determine forest parameters from optical satellite data. Trotter et al. (1997) found poor correlation between the SWIR band (TM band 5) and plantation forests, however, this was most likely due to the lack of complexity and shadows within the even-height plantation forests. The correlation between spectral data and wood volume tends to be stronger for younger stands than older stands. Horler and Ahern (1986) found Landsat’s two SWIR bands to be the most sensitive to forest vegetation density, particularly in the case of regenerating forest stands.

Clear-cuts are easily identifiable, with increased reflectance from the visible and SWIR bands and changes in the NIR reflectance (NIR may increase or decrease depending on the management activities and ground/field layer reflectance; Olsson, 2009). Determination of wood volume using Landsat and SPOT data has been hampered in several studies by the limited dynamic range of the spectral data, as reflectance of forest tends to be relatively low.

In an article by Nilson et al. (“Forest reflectance Modeling: Theoretical Aspects and Applications”, Ambio 32:535-541), results are presented from work with computer models of radiation in forest canopies, and verified with radiometer measurements in field and satellite data. The work has been carried out by a leading group at Tartu Observatory (partly in co-operation with SLU). According to the research presented in the article, the driving factors behind the reflectance from forest canopies are:

- canopy closure
- leaf area index
- tree species composition
- the understory vegetation

Unfortunately, the above factors are only partly correlated with forestry relevant information like stem volume, diameter and stem number. It is also stated in the article that the spectral signatures not changes much after 30-40 years.

Satellites which take frequent spectral images (e.g., an image every day, or every 5 days) create the opportunity to use phenology information to help identify different species. An example is that tree species which generally leaf-out earlier in the spring than others, can potentially be identified using this information. In other words, the difference in the development of vegetation types can give us yet another dimension of information to use in their identification.

### 3.7.2. Three-dimensional data sources

The most important data source to be introduced in the past decade for characterizing the forest has been three-dimensional data from airborne LiDAR. LiDAR allows a very accurate measurement of object locations (x,y) and height (z). For this reason, LiDAR is extremely useful for measuring of tree height, which in turn can be used in allometric equations.
for determination of other height related forest variables such as wood volume or stem diameter. Due to LiDAR’s ability to determine x,y position as well as penetrate the canopy, vegetation density can also be determined. The measure of vegetation density can be dependent on the number of laser hits per m² and also the characteristics of the vegetation.

LiDAR is currently often acquired using a laser in the near-infrared wavelength, and reflectance intensity is a parameter recorded in the returned data. However, due to a lack of information about the processing of these data (often due to company secrets), they often cannot be used to determine information about the vegetation types. Multi-spectral LiDAR are currently being introduced, and with further research, the multi-spectral information may be used to determine both forest structure characteristics accurately (height, density), as well as vegetation information (vegetation type).

It is not uncommon to combine the spectral information from satellite images with three-dimensional information from LiDAR, image-matching, or radar, in order to get the optimal information from each system.

3.8. The issue of scale
The importance of scale issues deserves special. Satellite data are often categorized by the pixel size (i.e., spatial resolution):

- Low or coarse resolution: 200-1000 m pixel size
- Medium or moderate resolution: 10-200 m,
- High or fine resolution: 1-10 m,
- Very high: < 1 m.

In the remote sensing context, Woodcock and Strahler³ use the term “spatial resolution” to refer to the sensors’ ability to resolve the spatial detail of the landscape. Strahler et al.⁴ described two different types of models representing the interaction between satellite spatial resolution and the scale of the objects being observed. These were \( H \)-resolution and \( L \)-resolution, in which \( H \)-resolution image pixels are smaller than the objects observed, and in \( L \)-resolution where the objects are smaller than the image pixels. As an example, for forests, \( H \)-resolution may translate to “several pixels per tree”, while \( L \)-resolution translates to “many trees per pixel.” Several studies have shown decreasing thematic classification accuracy with increasing spatial resolution.

In general, coarse resolution satellite data will have more pixels containing mixtures of cover types, making estimation of vegetation parameters more difficult. \( H \)-resolution imagery does not assure higher classification accuracies, as aggregated spectral information of the landscape may be necessary for accurate classification. For example, not only the spectral

³ Woodcock and Strahler, 1987
⁴ Strahler, 1986
reflectance from tree crowns but also the shadow cast by the tree crowns is an important source of information for the estimation of forest parameters. A moderate resolution pixel will capture both tree crown and its associated shadow. Higher spatial resolution data may provide more thematic detail, but the trade-off in pixel size is generally paid for with a smaller scene area coverage, resulting in a potentially more costly and complex mapping project.

The processing algorithms required to produce map information are likely to differ based on the spatial resolution of the data. For example, identifying tree cover with very high resolution data may require an aggregation of pixels into tree crowns (e.g., segmentation), while coarse resolution data require a breaking-down of the information in the pixel (e.g., spectral unmixing). The production of global land cover data often requires the use of coarse resolution data, which has stimulated the use of sub-pixel estimation methods to capture information about the heterogeneity within the larger pixel.

Different landscapes present different levels of heterogeneity and transitions between land cover types can be distinct or fuzzy, and hard to define even in the field. The spatial composition of the landscape structure, such as the forest stand sizes present, and the diversity within it such as presence of elements like bedrock outcrops, wetlands, water bodies, and roads, will have an effect on the result. The properties of the landscape and the goals of the mapping project exert important influences on the appropriate choice of remotely sensed data that have an “optimal spatial resolution”5. Some research has been done on determining “optimal pixel size”, while others have proposed using multiple-scale remotely sensed data for land cover classification.

5 Woodcock et al., 1988
4. OPTICAL SATELLITE DATA

**Ground Resolution Cell (GRC).** The size of the image grid cell, or pixel, which results from the sensor specifications.

**Thematic Mapper (TM).** The sensor aboard Landsat 4-7, which with its repeat coverage and 28.5 m ground resolution cell, made the optical satellite data from Landsat TM widely used.

When the term “optical” remote sensing data is used, it is meant that the sensor operates within the optical spectrum, of which the 400-780 nm wavelengths are visible to the human eye (e.g., blue, green, red). The spectral data recorded by the sensor is often related to the vegetation or other objects on the Earth’s surface. By analyzing the characteristics of the spectral reflectance, land cover types can be identified and mapped. The sensors used are most often placed in airplane or satellite platforms, and the images produced offer repeated views over the Earth’s surface.

4.1. Introduction to optical satellite data

Operational acquisition of optical satellite data has been taking place since the 1970s. Sensors often record information in the visible and near-infrared spectrum, and may include information in the short-wave infrared and thermal infrared spectrum. There are a number of different satellite data programs which provide data to the user community, and these are described in this chapter.

Countries often build their own sensors, where the characteristics of the sensors are often decided based on what is needed, what is possible, what has historically been used, costs, and market niches. Satellites may be state-owned and the data may be distributed free or at a cost, or corporately owned, where data are distributed often at a cost. The building of sensors and the cost of state-owned data has historically been politically influenced, yet there is currently a trend toward a free-data policy, as the use of data has been seen as beneficial to science and the economy. Since each satellite has the potential to view the whole globe, and since many countries want to have the technical capability to operate remote sensing satellites, but few countries are willing invest in redundant monitoring systems, there is a large need for international cooperation where satellite operators coordinate their plans. Organisations for such coordination are CEOS, and on a more general level also GEO.

4.1.1. Basic characteristics of optical data

The basic characteristics of remote sensing data have been named in Chapter 2, and for optical satellite data include spectral resolution, the number of wavelengths, spatial resolution, radiometric resolution, temporal
resolution, and image extent. Some properties of the satellites affect these characteristics, and are mentioned below.

**Satellite orbits**

Satellites have orbits that can vary in characteristics. They can have a polar or geo-stationary orbit, and they can have different heights above the earth’s surface.

*Polar orbit:* each orbit around the earth passes near the North Pole. Most earth resource satellites successively cover different parts of the earth for each orbit and repeat this pattern with a few weeks interval (the time before the same track is repeated was for example 16 days for Landsat 4, 5, and 7, and 21 days for the SPOT satellites). These satellite orbits are often *Sun synchronous* which means that the satellite moves in relation to the sun so that a given place is always recorded at the same time of the day.

*Geo-stationary orbit:* A very special case of satellite orbits is the geo-stationary orbit at about 36000 km exactly above the equator. A satellite placed in this circle around the earth will rotate with the same angle speed as the earth itself, and will then appear to be stable over one position above the equator. There is a system of five weather satellites in this geostationary orbit, each frequently viewing a complete view of the earth from one specific position. The European contribution to this constellation is called METEOSAT and the US is named GOES.

Most earth resource satellites, such as Landsat and SPOT, have a polar and sun-synchronous orbit, and have an altitude of about 700 – 900 km above the earth. However, there are also polar orbiting weather satellites, for example the NOAA series, the main difference being that the AVHRR sensor onboard the NOAA satellites have larger pixels and a wider swath. Each orbit around the earth only takes a few hours (for example 99 minutes in the case of Landsat 4-5).

There are also satellite orbits that rotates the earth on a relatively low altitude, without reaching the most northern or southern latitudes. One such example is the International Space Station (ISS). Thus, remote sensing sensors placed on ISS will not reach for example Sweden.

### 4.1.2. Common optical satellite data programs

#### 4.1.2.1. The Landsat program

**Landsat 1-3**

In the 1960s and 70s, for the persons that knew about military satellites, and for the larger number that had insights in the state of the art technology used to map the surface of the moon before the Apollo landings, it became more and more frustrating that this technology could not be used to map the resources of the earth as well.
To meet this criticism, the Landsat program was created. The Landsat 1 satellite was launched in 1972 (Table 4.1). It had a multispectral scanner (MSS) that produced digital images with 79 m pixels (often rounded up to 80m) in four spectral bands (nominally Green, Red, NIR1 and NIR 2).

The reason why the counting started on MSS 4 was that the first three bands were devoted to another sensor, a TV camera like system called Return Beam Videcon. These data are seldom used, whereas the MSS data can be used as an early reference for land cover change studies.

Landsat 1 was followed by the quite similar Landsat 2, launched 1975 and Landsat 3, launched 1978. MSS is still the best source for old satellite data before 1984.

**Landsat 4-8**

The US Landsat program became a success story, and Landsat 4 that was launched 1982, had an improved sensor named Thematic Mapper, with six reflective bands from blue to mid infrared, with 30 m pixels, and a Thermal band with 120 m pixels. The Thematic Mapper (TM) had better spatial resolution (30 m pixels) in six reflective bands, nominally in blue, green, red, NIR, SWIR1 and SWIR2. In addition, the TM sensor had a thermal band with 120 m pixel size: TM 6. The reason why the order between TM 7 and TM 6 doesn’t follow the order of the wavelength was that TM 7 was decided at a very late stage. This numbering has then been kept on the following TM sensors.

Landsat 4 soon got technical problems, but the similar Landsat 5 was launched 1984 and worked for more than 25 years, which is exceptional, since the design lifetime for a remote sensing satellite is normally about 5 years. Thus, Landsat 5 Thematic Mapper (TM) data is a very important source of old and current satellite data. Landsat 5 was collecting imagery into the year 2011 when it developed more problems, requiring that its operation be turned to an extremely minimum (and non-operational) level.

Landsat 6 was launched by a private company, but never reached orbit. It’s said to have crashed into the ocean after launch.

Landsat 7, launched 1999, was again a US government owned satellite. The sensor was now called ETM+. The main differences were a panchromatic band (TM 8) with 15 m pixel size, and thermal band (TM 6) with 60 m pixels. In May 2003 the Landsat ETM+ sensor developed a problem with its Scan Line Corrector, which resulted in images where the data had “line-drops”. Although there have been many attempts to correct these data and fill them in with data from adjacent images, Landsat 7 could not be seen as an operational instrument after 2003.

As of 2012, neither Landsat 7 nor Landsat 5 were able to produce usable data. This resulted in what researchers call “the Landsat data gap”, where the continuity of the Landsat program, operating since the 1970s, had been broken. The launch of Landsat-8 (also called the Landsat Data Continuity
Mission or LDCM) occurred in April 2013, solving this problem. Landsat-8 has a push-broom sensor called OLI (Operational Land Imager) but with specifications much like the previous Landsat TM sensors, with the main difference being higher radiometric resolution, and the addition of a coastal-blue and thermal wavelength bands.

Table 4.1. Satellites in the Landsat program

<table>
<thead>
<tr>
<th>Satellite</th>
<th>Launched</th>
<th>Decommissioned</th>
<th>Main sensor</th>
<th>Bands and Pixel sizes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Landsat 1</td>
<td>1972</td>
<td>1978</td>
<td>MSS</td>
<td>4 bands with 80 m pixels</td>
</tr>
<tr>
<td>Landsat 2</td>
<td>1975</td>
<td>1982</td>
<td>MSS</td>
<td></td>
</tr>
<tr>
<td>Landsat 3</td>
<td>1978</td>
<td>1983</td>
<td>MSS</td>
<td></td>
</tr>
<tr>
<td>Landsat 4</td>
<td>1982</td>
<td>2001 (last data 1993)</td>
<td>TM</td>
<td>6 bands with 30 m pixels + one 15 m panchromatic + one thermal band</td>
</tr>
<tr>
<td>Landsat 5</td>
<td>1984</td>
<td>last data 2011</td>
<td>TM</td>
<td></td>
</tr>
<tr>
<td>Landsat 6</td>
<td>1993</td>
<td>1993 (failed at launch)</td>
<td>ETM</td>
<td></td>
</tr>
<tr>
<td>Landsat 7</td>
<td>1999</td>
<td>sensor problem 2003</td>
<td>ETM+</td>
<td></td>
</tr>
<tr>
<td>Landsat 8</td>
<td>2013</td>
<td></td>
<td>OLI</td>
<td>8 bands with 30 m pixels + one 15 m panchromatic + two thermal bands with 100 m pixels</td>
</tr>
</tbody>
</table>

4.1.3. The SPOT program

The first satellite in the French SPOT satellite series, partly owned also by Sweden, was launched 1986. SPOT had a more modern push-broom scanner, based on a CCD-array, and better spatial resolution (10 m panchromatic pixels and 20 m colour). The SPOT program was an attempt to create a commercial market, and there have been seven SPOT systems launched. Data for the government owned SPOT 1-5 is summarized in Table 3.2. The SPOT satellites have now been taken over by the company Airbus and the recent SPOT 6 and 7 belongs to the very high resolution VHR category of commercially operated satellites and is therefore listed in Table 4.4 instead. The SPOT satellites have two push-broom sensors that are pointable sidewise. The original HRV sensors could deliver either a 20 m pixel IR colour image based on three spectral bands (Green, Red, NIR), or a panchromatic image with 10 m pixels. The HRVIR sensor onboard SPOT 4 had a mid-infrared band as well (which is good for forestry). The HRG sensors onboard had better geometric resolution (10 m colour, 5 m panchromatic). SPOT 5 also carries the HRS, a 3-line scanner for along track stereo.
Table 4.2. Satellites in the SPOT program

<table>
<thead>
<tr>
<th>Satellite</th>
<th>Launched</th>
<th>Main sensors</th>
<th>Bands and pixel sizes</th>
</tr>
</thead>
<tbody>
<tr>
<td>SPOT 1</td>
<td>1986</td>
<td>2 HRV</td>
<td>3 multispectral bands green - NIR with 20 m pixels or 1 panchromatic band with 10 m pixels</td>
</tr>
<tr>
<td>SPOT 2</td>
<td>1990</td>
<td>2 HRV</td>
<td></td>
</tr>
<tr>
<td>SPOT 3</td>
<td>1993</td>
<td>2 HRV</td>
<td></td>
</tr>
<tr>
<td>SPOT 4</td>
<td>1998</td>
<td>2 HRVIR</td>
<td>3 multispectral bands green – SWIR with 20 m pixels and the red band with 10 m pixels</td>
</tr>
<tr>
<td>SPOT 5</td>
<td>2002</td>
<td>2 HRG, +1 HRS for stereo images</td>
<td>3 multispectral bands green – NIR with 10 m pixels and one SWIR band with 20 m pixels. One panchromatic band with 2.5 – 5 m pixels</td>
</tr>
</tbody>
</table>

4.1.4. The Sentinel Program

The European Union has launched Copernicus, which is an ambitious program for earth observation. It consists of several series of earth observation satellites, as well as connection to in situ observations and operational services. The satellite series are called Sentinels, and are developed by the European Space Agency (ESA). Of most interest for forestry is data from Sentinel 2 which is a series of Landsat-like satellites. Sentinel 2a was launched 2015 and Sentinel 2b is planned for launch 2017.

The sensor has 13 bands with 10, 20, or 60 m pixel size, depending on wavelength (Table 3.3). The field of view is 290 km, which enables a Sentinel 2 satellite to pass a given point every 5 days, or even more frequently at high latitudes, like Sweden. The data are free and data over Sweden will be obtainable from a national archive. Thus, the Sentinel 2 data should be of high interest for forestry.
Table 4.3. Spectral bands provided by the Sentinel 2 satellites

<table>
<thead>
<tr>
<th>Sentinel-2 Bands</th>
<th>Central Wavelength (µm)</th>
<th>Resolution (m)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Band 1 - Coastal aerosol</td>
<td>0.443</td>
<td>60</td>
</tr>
<tr>
<td>Band 2 – Blue</td>
<td>0.490</td>
<td>10</td>
</tr>
<tr>
<td>Band 3 – Green</td>
<td>0.560</td>
<td>10</td>
</tr>
<tr>
<td>Band 4 – Red</td>
<td>0.665</td>
<td>10</td>
</tr>
<tr>
<td>Band 5 - Vegetation Red Edge</td>
<td>0.705</td>
<td>20</td>
</tr>
<tr>
<td>Band 6 - Vegetation Red Edge</td>
<td>0.740</td>
<td>20</td>
</tr>
<tr>
<td>Band 7 - Vegetation Red Edge</td>
<td>0.783</td>
<td>20</td>
</tr>
<tr>
<td>Band 8 – NIR</td>
<td>0.842</td>
<td>10</td>
</tr>
<tr>
<td>Band 8A - Vegetation Red Edge</td>
<td>0.865</td>
<td>20</td>
</tr>
<tr>
<td>Band 9 - Water vapour</td>
<td>0.945</td>
<td>60</td>
</tr>
<tr>
<td>Band 10 - SWIR – Cirrus</td>
<td>1.375</td>
<td>60</td>
</tr>
<tr>
<td>Band 11 – SWIR</td>
<td>1.610</td>
<td>20</td>
</tr>
<tr>
<td>Band 12 – SWIR</td>
<td>2.190</td>
<td>20</td>
</tr>
</tbody>
</table>

4.1.5. Very high resolution satellites

Remote sensing satellites with sensors that produce images with much better spatial resolution than Landsat, and later SPOT, were from about 1960 to the beginning of the 1990’s only available for military intelligence. This changed however with the end of the Cold War and the collapse of the Soviet Union. Russian space authorities then started to sell spy satellite images on the open market. The pixel size had been degraded to 2 m, but this was still much better than any other satellite images available on the open market at that time.

The USA responded to this in two ways. By releasing their old CORONA images for sale on internet, and by licensing out the right to build “civilian spy satellites” to private companies. The first of these very high resolution satellites was IKONOS, that was launched by Lockheed 1999 and produced panchromatic images with 1 m pixel size and IR colour images with 4 m pixels. Later in 2001 followed Quick Bird that produced images with 0.5 pixel size in the panchromatic band. There has since been many follow on satellites of similar type.

Today, there are many nations that have launched, or are planning to launch, remote sensing satellites which produces images with pixel sizes in the order of 1m. Examples are France, South Africa, Israel, Korea, Taiwan and India. Governments are willing to invest in this technology, much for military reasons, but since it is not secret any more, they are often also willing to sell images on the open market as well, this strategy is called dual use. Thus, the “very high resolution segment” can be regarded as
stable. Table 4.4 lists some of the more important high resolution satellite sensors.

These very high resolution satellites produce imagery that is almost like orthophotos from aerial cameras. Each image also covers however a very limited area (often in the order of $20 \times 20$ km). Thus, they compete mostly with aerial photos and it is difficult to obtain cloud free data for large areas like whole countries. The market for very high resolution satellites is probably limited in countries with a regular supply with governments subsidized air photos, such as Sweden.

One of the latest developments is coming from PlanetLabs, which is sending up “shoebox satellites”, which are very small, inexpensive satellites. PlanetLabs aims to send up 100-200 shoebox satellites so that they can achieve high temporal resolution, namely to cover every spot on the Earth, every day. The data, provided by a commercial company, are not free. We can expect to see more of this type of development in the future.

Table 4.4. Some very high resolution satellite programs that produces imagery for the civilian market.

<table>
<thead>
<tr>
<th>Satellite</th>
<th>Year Launched</th>
<th>Multispectral bands</th>
<th>Multispec bands pixel size</th>
<th>Pan band pixel size</th>
<th>Image Swath</th>
</tr>
</thead>
<tbody>
<tr>
<td>IKONOS</td>
<td>1999</td>
<td>B, G, R, NIR</td>
<td>3.2 m</td>
<td>0.8 m</td>
<td>11.3 km</td>
</tr>
<tr>
<td>QuickBird</td>
<td>2001</td>
<td>B, G, R, NIR</td>
<td>2.4 m</td>
<td>0.6 m</td>
<td>16.5 km</td>
</tr>
<tr>
<td>WorldView-1</td>
<td>2007</td>
<td>-</td>
<td></td>
<td>0.46 m</td>
<td>17.6 km</td>
</tr>
<tr>
<td>WorldView-2</td>
<td>2009</td>
<td>B, G, R, NIR + 4 more</td>
<td>1.84 m</td>
<td>0.46 m</td>
<td>16.4 km</td>
</tr>
<tr>
<td>WorldView-3</td>
<td>2014</td>
<td>28 bands 400-2365nm</td>
<td>1.24-30 m</td>
<td>0.31 m</td>
<td>13.1 km</td>
</tr>
<tr>
<td>WorldView-4</td>
<td>2016</td>
<td>B, G, R, NIR</td>
<td>1.24 m</td>
<td>0.31 m</td>
<td>13.1 km</td>
</tr>
<tr>
<td>GeoEye-1</td>
<td>2008</td>
<td>B, G, R, NIR</td>
<td>1.65 m</td>
<td>0.4 m</td>
<td>15.2 km</td>
</tr>
<tr>
<td>RapidEye (5 satellites)</td>
<td>2008</td>
<td>B, G, R, NIR + 1 “red edge”</td>
<td>6.5 m</td>
<td>-</td>
<td>77 km</td>
</tr>
<tr>
<td>Pleiades 1A</td>
<td>2011</td>
<td>B, G, R, NIR</td>
<td>2 m</td>
<td>0.5 m</td>
<td>20 km</td>
</tr>
<tr>
<td>Pleiades 1B</td>
<td>2012</td>
<td>B, G, R, NIR</td>
<td>2 m</td>
<td>0.5 m</td>
<td>20 km</td>
</tr>
<tr>
<td>SPOT 6</td>
<td>2012</td>
<td>B, G, R, NIR 6 m</td>
<td>1.5 m</td>
<td></td>
<td>60 km</td>
</tr>
<tr>
<td>SPOT 7</td>
<td>2014</td>
<td>B, G, R, NIR 6 m</td>
<td>1.5 m</td>
<td></td>
<td>60 km</td>
</tr>
</tbody>
</table>

How many satellites are there in orbit?

There are all types of satellites, some of them used for communication, and some for Earth Observation (Fig 4.1). There are approximately XX Earth Observation satellites now in orbit.
4.1.6. **Satellite scenes and image extents**

Satellite data are traditionally distributed as square scenes. Most satellite systems have a *World Reference System* (WRS) that assigns a unique number to each potential satellite scene, often based on a Path (north to south) and Row (east to west) number. For example, the Landsat image center closest to Umeå is from Path 193 and Row 15, and is referred to as 193/15. Landsat images have a fixed WRS identifier (the same area will always have the same WRS number). In contrast, SPOT images do not have a constantly fixed area, and for this reason, the WRS number is approximate. To uniquely identify a satellite image, you need the WRS number, as well the date the scene is received. (For satellites that have more than one sensor, such as SPOT, you might also need the name of the sensor). Often, the WRS identifier makes up the file name of the satellite image.

In the case of Landsat OLI, a scene is $185 \times 185$ km, and in the case of SPOT $60 \times 60$ km. Theses “full scenes” covers the whole field of view (FOV) sensed by the satellite sensor in the east-west direction. However, in the north-south direction, the image received at the satellite station is often from a technical point of view, a many scenes long strip. Thus, it is sometimes possible to order a “non-standard” scene where the limit for the scene is adjusted north-south according to the customer order.

Many satellites, like e.g. SPOT, have *pointable sensors*, making it possible to increase the chances to get cloud free images over a certain area. The use of oblique viewing will however be at the cost of not viewing the area under the satellite. If it is important to obtain images over a certain area a certain year with such a sensor, it is therefore often possible, and advisable, to order a *programming* of the satellite in advance. Such a programming should preferably be ordered a few months in advance and might cost extra.

The development today is that most users of satellite data will use ready-made seamless mosaics, and not handle the raw-data in the individual scenes themselves. A very popular example of this is Google Earth. For
professional use, there is however most often a need to not only be able to view the image data, but also know the sensor characteristics and the date and processing for the image.

**Downlinking of data**

Getting the huge amounts of image data down to the earth has always been a bottleneck for satellite remote sensing. The early spy satellites used film capsules sent down with parachutes. Later earth resource satellites with digital sensors (such as the early Landsat satellites) downlinked the data to a few receiving stations with large parabolic antennas. For areas outside these receiving stations, images could be tape recorded and downlinked when a receiving station was passed. The network of receiving stations has successively become denser and denser, but onboard recording of data is still important. The sensitive tape recorders are now replaced by solid state computer memory. For downlinking of data and also for sending control commands to the polar-orbiting satellite, it is an advantage to have receiving stations with a northern location. Two such stations are Esrange in Kiruna and the Norwegian station at Svalbard which in 2010 had 18 parabolic antennas for receiving satellite data.

Some remote sensing satellites, for example Landsat 4, have also had the possibility to send the data to a central receiving station that might be on the other side of the earth, by using a telecommunication satellite in geostationary orbit as a link.

**4.2. Basics in image interpretation**

In the following section, we give a brief overview of the process of analyzing optical satellite data to create information on forests and other land cover. More detail about the analysis methods is given in Chapter X.

When beginning a project, there can be a number of steps involved, which we have summarized as the following:

- Knowing what you need or want for your final product;
- Determine data sources (remotely sensed data as well as reference data);
- Pre-processing of remotely sensed data;
- Decide analysis method;
- Analysis;
- Initial control of the map product;
- Post-processing of product; and,
- Final accuracy assessment
4.2.1. Knowing what you need or want

The desired final goal of the project is often known. External data users may order a product, and turn to remote sensing experts to deliver this. It is up to the remote sensing expert to know whether the data at hand can meet the goals and costs of the project. The first step is to identify the goals, the potential data sources, the time and cost, and estimate the potential accuracy (based on previous studies) of the outcome.

If you want to create a thematic map, with a number of land cover classes, then you will need to develop a classification scheme. A classification scheme consists of the class names and class definitions. The classification scheme should cover all possible cases and be mutually exclusive so that every case should belong to one and only one class.

The project may develop its own classification scheme, or use existing ones. Examples of existing ones are the FAO classification system, the Anderson system, or the ISO system, among others. It is often useful if classification schemes are hierarchical, and depending on outcomes, may be collapsed into broader classes if necessary.

It is common that a classification scheme consists of either Land Cover classes, or a mix of Land Cover and Land Use classes. Land cover refers to the actual cover on the ground, while Land Use refers to the use. An example is “Wheat” would be the Land Cover class, while “Agriculture” is the Land Use class. It is often more difficult to determine what the use for an area is as opposed to the cover.

It is also of importance to know what your Minimum Mapping Unit (MMU) in your final map product will be. The MMU is the smallest area of a class that you will map. The MMU is dependent on the original pixel size of your data, and the ability to detect this phenomenon. For example, if you state that your final map MMU is 0.5 ha, then you will need to generalize your final map product to areas of this size. Any landscape object smaller in size than your decided MMU will be assimilated or ignored. The MMU and the properties of the landscape can affect the reference data collection, the post-processing, and the accuracy assessment numbers.

4.2.2. Determine the data sources

Different remote sensing data are suitable to different tasks. A global coverage of forest? It will be important to use data with large scene extent, and global coverage. A smaller area mapping of urban expansion? A long term availability of higher resolution data may be most suitable. A map over a rainforest area? Here it is not enough to know that a sensor exists, but whether usable data really exist over the area to be mapped.

Often, a bigger challenge is to obtain the reference data needed for both interpretation and accuracy assessment of a mapping project. This is particularly true if the remote sensing data were acquired some time ago (i.e., earlier in time than the current date). Refer to the section on Reference Data to explore this issue more.
4.2.3. Decide on the analysis method and analysis
There are many different methods used to process the remotely sensed data into map data. The method chosen often depends on the input data available, and the desired result. Some analysis methods require more reference data than others, some analysis methods are used only for creating thematic maps and others used only for creating continuous value maps. Some analysis methods must be chosen based on the characteristics of the data (e.g., coarse resolution vs high resolution data). Chapter X goes into more detail about the characteristics and uses of different analysis methods.

4.2.4. Reference data pre-processing
Even reference data themselves require a pre-processing step, which might consist of controlling the quality of the reference data, forecasting or back-casting the reference data, determining the suitability, removing outliers, and perhaps adding more reference data if deemed necessary.

4.2.5. Pre-processing of optical satellite data
The radiance recorded by a satellite sensor is affected by several factors, including the atmosphere, topography, vegetation composition, solar illumination angles, and sensor characteristics. Optical satellite data have characteristics that require some pre-processing steps which other remotely sensed data do not. For example, due to the visible wavelengths that may be influenced by atmospheric haze, there may be a need to reduce atmospheric effects. Some of the common pre-processing steps, namely atmospheric correction and topographic correction, are mentioned below. The need of doing these steps depends on the goals of the project, as well as time and cost limitations.

4.2.5.1. Factors affecting the DN values
The way we use optical satellite data in forestry is most often to compare the digital numbers (DN) to field reference data, in order to convert the DN values from a combination of spectral bands to either discrete classes, or to forest data with continuous values, e.g. stem volume. The DNs are the values delivered in the satellite image product, and are related to the true reflectance from the objects plus including the influence of atmospheric attenuation (as explained in Chapter 2). There are computer programs for converting the DN values for each spectral bands to units of reflectance. In practice, this will be a linear transformation of the type:

\[ \text{Reflectance} = a + b \times \text{DN} \]

One advantage with this procedure is that reflectance is a physical unit that is more comparable between sensors, and with data in the literature than the DN values. However, since the atmospheric correction is just a linear transformation, it will not improve estimates of forest variables or classifications into discrete classes, when they are made by the aid of ground reference plots.
It is also difficult to replace the need for ground reference plots with a library with reflectance values. One reason for this is that the estimation of the coefficients $a$ and $b$ in the above function seldom is accurate enough. The coefficients depend both on the atmospheric optical thickness on that actual day, and on the calibration of the sensor and both of these factors vary over time and are difficult to estimate accurate enough for the purpose of creating general relationships for forest estimation.

Furthermore, the reflectance from a forested pixel depends not only on the size and amount of trees, but also on the length of the shadows, and the status of the field layer vegetation. Both these factors vary considerably with the season of the year, which also complicates the use of reflectance as a general measure for forest status.

The above mentioned factors: atmosphere, sensor calibration, shadow length, solar angles and vegetation season, will also influence both scenes in a bi-temporal change detection. Thus, the best way to find changed areas is to normalize the DN values in both scenes to each other with statistical techniques rather than using physically based atmospheric correction.

The atmospheric optical thickness might vary over different parts of a satellite scene. This might be studied by contrast stretching, using the most shortwave of the available bands, e.g. the blue band. It is best to totally avoid using scenes with visible haze for computer based analysis. Furthermore, it is always a danger to include the blue band in automated analysis for this reason. The longer wavelength in the range blue – SWIR, the less atmospheric influence, thus, automated analysis over large areas should preferably be based on long wavelengths, and the blue, and even the green bands should be avoided if possible.

The topographic characteristics of an area, such as slope and aspect, in combination with the angles of the sun’s position (typically the solar zenith and azimuth angles) result in illumination differences within a satellite image. Topographic correction is used to directly manipulated the DNs in the satellite image so that a vegetation class will have similar spectral response whether on a north facing slope (facing away from the sun) or on a south facing slope (towards the sun).

4.2.5.2. Topographic normalization, C-correction

Topographic normalization methods are often grouped as either photometric/photometric-empirical which includes the semi-empirical corrections such as Minnaert correction\(^1\), $b$-correction\(^2\), and C-correction\(^3\), or physically-based models\(^4\). Gu and Gillespie (1998) suggested using different topographic normalization methods for forested vegetation, namely the more forest-appropriate Sun-Canopy-Sensor (SCS) correction.

---

\(^1\) Smith, Lin, and Ranson 1980
\(^2\) Vincini and Frazzi 2003
\(^3\) Teillet, Guindon, and Goodenough 1982
\(^4\) Shepherd and Dymond 2003, Soenen et al. 2008
While for non-forested areas the C-correction or statistical-empirical correction is most appropriate. The SCS model was later modified by Soenen et al. (2005) to include the c-parameter, called the SCS+C method.

A relatively simple method for topographic normalization is the C-correction. This example may help understand the process of topographic normalization. The C-correction was developed to perform topographic correction for non-Lambertian reflectance behavior, as the cosine correction was suitable for Lambertian reflectors only\(^5\). It is based on the linear relationship between the spectral reflectance recorded for a pixel and the corresponding cosine of the solar illumination incidence angle, \(i\). The cosine of \(i\) can be calculated as a function of the local terrain slope and aspect, as well as the solar illumination angles upon the surface at the time of satellite data acquisition (Eq. 1).

\[
\cos i = \cos z \cdot \cos s + \sin z \cdot \sin s \cdot \cos (\Phi_a - \Phi_n) \tag{1}
\]

where \(i\) is the solar illumination incidence angle with respect to surface normal, \(z\) is the solar zenith angle, \(s\) is the terrain slope angle, \(\Phi_a\) is the solar azimuth angle, and \(\Phi_n\) is the terrain aspect angle (Fig. X).

Linear regression with cosine of \(i\) as the independent variable and reflectance (\(\hat{\rho}_\lambda\) is the topographically-influenced (\(t\) reflectance of band \(\lambda\)) as the dependent variable (Eq. 2) is used to estimate intercept \((b)\) and slope \((m)\).

\[
\hat{\rho}_\lambda = b + m \cdot \cos i \tag{2}
\]

The \(c\)-parameter is calculated as \(b\) divided by \(m\) (Eq. 3). The relationship between reflectance and cosine of \(i\) is wavelength dependent (Teillet, Guindon, and Goodenough 1982), therefore, a \(c\)-parameter is calculated for each wavelength band.

\[
c_\lambda = \frac{b}{m}, \tag{3}
\]

The \(c\)-parameter is then added to the numerator and denominator of the cosine correction to form the C-correction equation (Eq. 4).

\[
\hat{\rho}_{\lambda h} = \hat{\rho}_\lambda \frac{\cos \theta_z + c_\lambda}{\cos \theta_z + c_\lambda}, \tag{4}
\]

where \(\hat{\rho}_{\lambda h}\) is the topographically normalized reflectance (\(h\) indicating “horizontal”) for band \(\lambda\), and \(c_\lambda\) being the \(c\)-parameter for band \(\lambda\). According to Teillet et al. (1982) and Meyer et al. (1993), the \(c\)-parameter is used as an approximation of diffuse sky irradiance.

\(^5\) Teillet et al. 1982
4.2.5.3. Image manipulation
Filters and indices are examples of manipulation to images. This text has not yet been written, and therefore we refer to the supplemental material for reading in the course.

4.2.6. Analysis
See Chapter 8 for more details on the analysis phase of an image processing project.

4.2.7. Initial control of the map product
Once you have run the first iteration of your classification, you may be excited to check the result. It’s likely though that the first run of your classification will not be the last. You will want to check your work by comparing the map product to some reference data anchored in reality. This may come from existing reference data, and/or that the remote sensing analyst goes out in the field. You may find that a certain class is poorly classified, and you need to adjust your input data or method to obtain the end result which meets the requirements of your project. The value of the remote sensing analyst visiting the study area and observing the actual vegetation and input data cannot be underestimated, as a deeper understanding of the relationship between the vegetation and the remote sensing data will only improve the outcome of the mapping process.

4.2.8. Post-processing map manipulation
Post-classification smoothing where isolated pixels are re-assigned with the class label of nearby pixels, with the aim of creating a “smoother” looking map. The result from supervised classification may require smoothing because it often looks more “pixelly” or has a “salt-and-pepper” look than does the outcome from unsupervised classification.

Different algorithms can be used for smoothing, such as a majority filter (also called a mode filter). There are also “eliminate and fill” algorithms. The window filters, such as a majority filter work like this:

- move a 3 by 3 pixel “filter box” over the classified image,
  for each pixel position, replace the center image in the 3 * 3 box with the most usual pixel value in the box.

The smoothing effect will be stronger if a larger filter, e.g. 5 by 5 pixels are used. Test what happens with the coding of the below classified image when a 3 * 3 pixel is moved over the image and the centre pixel is for each position of the box replaced with the most frequent value in the box.
4.3. Access to optical satellite data

The file size of satellite images is currently increasing, for example, the original Sentinel-2 images as a whole are as much as 6 Gb in size. However, the remote sensing community is attempting to make as many images as possible available from the archives. The access of optical remote sensing data depends on the sensor. Those that are freely available can be found from several websites, which are given in the following text. It is currently common that there will be several websites offering different processing or different availability of satellite data. For this reason, it is beneficial to look at more than one website. For commercially available satellite data, websites might also be used if satellite images have been archived by the company, or they may need to be contacted directly in order to program the desired image, which will be sent most likely via internet. Some of the most common access points for satellite data are SACCESS, USGS Earth Explorer or GLOVIS, Copernicus, the Global Land Cover Facility at University of Maryland, the European Space Agency, and the SPOT Sirius catalog. Links to these are given in the chapter on Internet links for this chapter.

4.4. The future of optical satellite data remote sensing

There are plans to send up a Landsat 9 OLI sensor in the year 2020. It is another step in the continuation of the long Landsat series. The Landsat series aims to keep the basic characteristics of the sensor through time, in order to have long time series of comparable data. The funding of the US Landsat program is, however, highly dependent on political decisions, and is often subject to fluctuations. Google Earth works closely with Digital Globe Inc. which has the WorldView satellite series, and has plans for future WorldView satellites. There is also a trend to commercialization of the space industry, and from this, PlanetLabs has started sending up “showbox” satellites. A good source of information on future satellite launches is provided in the internet links, and can often be found on the internet.
The European Space Agency plans to continue with the Sentinel space program, sending up new versions of Sentinel-2 satellites in the next decade. Other nations that have strong space programs with future plans for optical satellite data are India, with its ResourceSAT satellite series, and China.
AERIAL PHOTOGRAPHY: IMAGE INTERPRETATION AND DIGITAL PHOTOMETRY

Aerial photographs for forest remote sensing

After the Second World War the use of aerial photographs in Swedish forestry took off. Mainly images were used to delineate land use classes and forest or the so-called compartments of forest inventories and the planning of treatments. This was a major step forward because in the past much time had been spent to map compartments using only field methods.

Over the years, special techniques developed within the forest aerial photography technology. Digital images have replaced the analog, which also resulted in a shift of equipment and also created the basis for new methods. For a long time, the most common image material was black and white images (usually panchromatic) from funded campaigns, the so-called repeated photographing. These normally were taken from two altitudes: 4600 m (normal height) and 9200 m (altitude adjustments). For a brief period there were also pictures at 13 200 m (high altitude). The negative scales for these images was about 1:30 000, 1:60 000 (camera constant 15 cm) and 1:150 000 (camera constant 9 cm). Often stereo mounted hard copies of the images were used for field work, such as for forestry planning (figur 5.1).

Figur 5.1. equipment for the viewing of stereo images in the field. The aerial photos are taped to a picture holder and a so-called lens stereoscopic (Zeiss) is permanently attached with a string. Leg construction (Simon's model) allows one to unfold one image which is necessary to consider the whole stereo model.
Detail reproduction is of course worse with smaller-scale photography. This means that images taken from 4600 m height were dominant for forest image interpretation. Pictures from the higher altitudes were mostly used for orthophoto-production and easy updating of maps.

From the year 2005, the National Land Survey was shooting with the digital camera “Zeiss / Intergraph Digital Mapping Camera (Z / I DMC)” at a flying height of 4800 m. This gave pictures with the scale of 1:40 000 (camera constant 12 cm) and ground resolution cell about 48 cm, which is about the same resolution as in the analog photograph images taken at a flying height of 4600 m. (Note that in the north of the Sweden, forestry stakeholders financed photography to be carried out using color infrared film, which meant that the transition to digital technology took place one year later.)

Images recorded with the Z / I DMC and ground resolution cell of about 48 cm was the dominant image type between 2006 and 2012, but the other camera types with slightly higher resolution had been developed. Currently Vexcel Ultracam Eagle is the most common camera used by Lantmäteriet. Usually the flying altitude is selected so as to have a ground resolution cell of about 24 cm or about 48 cm.

Need of timeliness and better detail reproduction may warrant special photography from lower flying altitudes. The National Forestry Board acquired aerial photographs in the 1980s, with most at 3000 m altitude in order to make a comprehensive forest inventory (OSI). They then used the contact copies in approximate scale of 1:20 000.

For some special purposes also other image types have been used. In the 1980s, increased demands for treatments after harvesting were called for. This meant that some forestry companies carried out the so-called "Low altitude photography" of clearcuts. Typically, conventional color film and a medium format camera was used (6x6 cm image format). The advantage of the format was that cheaper cameras and small aircraft could be used. The downside was that the images covered smaller areas, which was ok when smaller areas were to be photographed, but not for example, forestry planning of large estates.

For some purposes, images taken with so-called UAV (Unmanned Aerial Vehicle) or drones can be useful. The Swedish National Forestry Board has its own equipment and there are also consulting firms that offer services including orthophoto production.

IR color images in Sweden have mainly been used for image interpretation in connection with the preparation of vegetation maps. Image scales has then been between 1:30 000 and 1:60 000. Almost all of Sweden was photographed with analog infrared color images over a period ranging between the mid 1970s and early 1990s.
In the forestry context, the infrared color images have been used, among other things, for the interpretation of forest damage and young forest inventory related to special surveys and experiments. The approximate photography scales has then usually been between 1: 2000 and 1:10 000. IR color images of normal height and high altitudes have been used in inventory of hardwood forests and determination of the forestry border in the mountain regions.

In the early 2000s in Sweden, more and more of the state funded analog photography programs were carried out with infrared color film, and in the forest counties large areas were partly financed by forest stakeholders. This meant that this type of image also became common in forest management planning and also for the production of digital orthophotos.

The more advanced digital aerial camera can register multiple wavelength bands than what was possible when shooting with film. This means that from the same image taken with a digital camera, both true color and infrared color images could be produced. When ordering digitally photographed images from the National Land Survey one can therefore select the type of picture which best suits the goal.

In more advanced image interpretation of analog images previously used diapositive interpreted either in zoom stereo microscope or the so-called analog or analytical stereo instruments (Figure 5.2 - 5.3) These types of instruments cannot be used to view digital pictures and they have been replaced by digital photogrammetric workstations, also called digital stereo instruments (Figure 5.4).
Figur 5.3. Analytiskt stereoinstrument Zeiss Planicomp P3. I de analytiska stereoinstrumenten återskapas en geometriskt riktig modell av verkligheten med matematiska metoder (till skillnad mot i de analoga instrumenten där man mäter i en analog modell av terrängen). Planicomp P3 var tidigare ett vanligt instrument bl.a. hos Lantmäteriet vid karteringen av Fastighetskartan. (Zeiss)

Figur 5.4. Digital photogrammetric workstation, or as it is also called, digital stereo instruments. Stereo model is considered in this case with the aid of polarimetric on the right screen. Map database presented on the left screen. The digital instruments are basically the same way that the analytical mechanics but has been replaced by electronics, the optics of the displays and the analog images of digital.

For advanced image interpretation of digital images requires high viewing equipment and software that allows stereo viewing, which is met in the digital stereo instruments.

The aerial photos should always be supplemented with information on timing of photography (especially the date). It is also of value with the data on
the general image data, for example, altitude, resolution and filtering. If the image is used for measurements, it is necessary with data on the camera (e.g., the camera constants).

**Introduction to aerial photography**

Aerial photography to be acquired for the purpose of mapping and image interpretation is carried out in straight flight paths. For the images to be viewed in stereo the photography usually must be done so that there is about 60% overlap between images. If larger areas are to be photographed there must be multiple parallel flight paths used. The paths are flown then to obtain a covering also between paths, usually about 30% (see also Figures 5.5 and 5.6).

![Aerial photography diagram](image.png)

**Figur 5.5**. Aerial photography is done in straight paths, usually with about 60% overlap between images.

![Aerial photography diagram](image.png)

**Figur 5.6**. Aerial photography is done in straight paths usually with about 60% overlap between images in the photo strip and about 30% overlap between the routes.
Altitude largely determines the measurement accuracy that can be obtained and also has a crucial importance for what is possible to identify and interpret the images. For economic reasons, choose most often as high altitude as possible with respect to the accuracy requirements. At moderate altitudes usually the rule of thumb that the uncertainty in the plan for well-defined objects are of the same magnitude as the digital images ground resolution. Measurement uncertainty in height is about 1.5 times larger.

The altitude is therefore chosen so that the images ground resolution at least equal to the requirement of uncertainty in the plan (Note that high ground resolution = low amount). While the requirement of the small details that can be identified are important for the selection of flight altitude, i.e., one must also take into account the type of mapping or image interpretation to be performed.

Flight path planning is for the most part done to cover the area to be photographed with as few paths and images as possible. For special photo acquisitions, it is possible to select how the paths placement can meet this requirement, which is usually called free flight path planning (figure 5.7). Exceptions to this may be that you want photographs for a particular map sheet, i.e., images covering map sheets sideways. This is called bound path planning and was especially common in the past. For example, Lantmäteriet acquisitions are usually planned so that the images cover a predefined map sheet (5x5 km) by some margin. This also means that north-south paths were normally photographed and had the turn carried out in the middle of map sheets (Figure 5.8). Bound path planning facilitated the subsequent work, including the preparation of orthophotos (see below), but is less important with the current digital technology.
Cameras

In aerial photography for measurement purposes, we usually use a so-called digital mapping camera. These have known and stable internal geometry. The internal geometry is measured (calibrated) by the camera manufacturer and is reported in a calibration protocol. The data measured is the location for the so-called image principal point H (see figure 5.9) and camera constant c, i.e., the distance between the centers of projection and image principal point. Additionally measured is the optical lens displacement. Displacement occurs when a light beam changes direction when it passes the projection center. This data is called the camera interior orientation data and then used in the photogrammetric processes. Figure 5.9 is a symbolic digital mapping camera with interior orientation data reported.
Earlier images were photographed with film but now use a CCD (Charge-Coupled Device). The most common type consists of a matrix of cells (CCD matrix) which are connected and which after exposure are electrically charged at different rates depending on the amount of light that hits each cell. In such a camera, the image has a central projection in a similar manner as in the previously used analog cameras (figure 5.10).

There are also sensors which consist of a single row of CCDs (CCD line). The advantage of these is that they are easier to manufacture than CCD arrays. With such a camera you have to continuously scan the landscape during the flight, and the result is a cylindrical projection (Figure 5.11). It also places high demands on the navigation equipment in the plane so that it continuously records the camera's position (in all directions) because each scan line must be connected to the previous and subsequent rows. This type of sensor has so far had little use in Sweden and therefore what is described in the text is primarily images from the CCD arrays.
It is difficult to manufacture a CCD array of sufficient size to be able to record images in large formats and with enough resolution for image interpretation and Aerial Photogrammetry. This has led to some camera manufacturers solving the problem by building sensors that consists of several cameras mounted next to each other where the cameras are slightly slanted and thereby depicts a larger ground area. "Side-looking images" from the different cameras rectified after shooting to nadir images and converted to a single central projection (Figure 5.12). In connection with this calculation corrects images for radial displacement and systematic errors and the end result is a nearly faultless central projection.

DMC cameras, which were the most common digital mapping camera in Sweden before 2013, and even Vexcel Ultracam Eagle used by the National Land Survey now, all consist of camera packages with multiple cameras. They have all four cameras that record high-resolution black and white (panchromatic) images in the manner described above. They also have four color cameras with lower resolution, which records either blue (B), green (G), red (R) and infrared (IR, so-called near infrared). Using a process called pan sharpening the low-resolution color images are combined together with high-resolution monochrome image and the final result will be either a natural color image or an infrared color image. The pan sharpened picture is usually perceived as a high-resolution color image at "normal" inzooming.

In the digital mapping camera fiducial marks are not required, location of the principal point is defined by the row and column values in the CCD array of the pixel corresponding to the image principal point.
Table 5.1 data are available for some of the digital flygmätkameror available on the market at present (2016).

<table>
<thead>
<tr>
<th></th>
<th>Z/I DMC</th>
<th>Vexcel UltraCamXp</th>
<th>Vexcel UltraCamE</th>
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<tr>
<td>Bildstorlek (MP)</td>
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<td>196</td>
<td>260</td>
</tr>
</tbody>
</table>

**Lantmäteriets flygfotografering**

The National Land Survey (Lantmäteriet) is responsible for the state funded aerial photography program in Sweden. Photographings on a large scale began after a parliamentary decision in 1937 when it was decided that a new economic map would be produced that was based on photos. Forestry was one of the driving forces to make this happen. The national photography program is typically carried out entirely by the Lantmäteriet, or earlier by ORDNANCE SURVEY, but nowadays about 40% of the photography was carried out by contractors.

Presently, LM takes aerial photographs over Sweden on average every three years. The interval between the photography per area varies between 2-10 years. Southern parts of the country and the Norrland coast is planned photographed every other year, with inland Norrland every 4th to 6th year and the mountains every sixth to tenth years (Figure 5.13).

The images taken every two years are planned to be carried out with about 24 cm ground resolution, alternately before and after leaf-out. The images photographed at longer intervals should be taken approximately 48 cm resolution and after leaf-out. The photos can be ordered either as orthophotos or stereo images with the orientation data for use in digital photogrammetric workstations.
Figur 5.13. Lantmäteriets plan for aerial photogramy. The green area, to be photographed with the two-year intervals, are scheduled to be taken both before and after the time of leaf-out (changing between each acquisition). Other areas will normally be photographed always after leaf-out.

**Ortofoton och fotokartor**


I en flygbild som är fotograferad med en kamera som ger en centralprojektion, förskjuts höga objekt, t.ex. berg eller trädkronor, ut från bildcentrum.
In the production of an orthophoto, a model of the ground height is used to control a recalculation of the image so that the map becomes a real orthogonal projection. Because it is the ground height that is used, then the tree crowns and rooftops will still be slightly displaced from the image center. If you want to compensate even for this effect, you need instead of a ground model to use a surface model, showing the altitude of the highest objects in each image pixel. When using a surface model to get a product called "true orthophoto". To produce a true orthophoto more images are required, namely to cover larger areas between the images, otherwise you have "holes" in the orthophoto where the image data is missing because the ground is obscured due to shadowing effect (so-called shadow projection), see Figure 5.14a-b.

5.14a In a normal orthofoto is fortfarande hustaken lägesförskjutna på hus som ligger i kanten den bild som ortofotot är framställt från. Foto: Lantmäteriet.

5.14b In a true orthophoto has been compensated even for displacements of protruding objects such as rooftops. This requires that more images are used, otherwise there will be holes in the image where the image information is taken. For example, the surface information is lacking due to shadow (see arrow) as compared to the image to the left.

**Determining wood volume with the LMV method**

In many contexts, we are interested in how much timber is available in an area, that is, the stock of wood or timber volume. Stand or compartmental estimates is a common method and for this purpose aerial photographs have been used to aid in various ways. Delineation of compartments are an application where aerial photographs have been most important. Often a preliminary delineation is done using aerial images which are then checked in the field and adjusted if necessary. Earlier mostly a simple stereoscope was used for this purpose (Figure 5.1).

A common method has been to use stereo images in the field to estimate volume with subjective methods. Often for reasons of time, measurements of basal area could not be made because not enough areas could be assessed within the stand. For this reason, aerial images were used to assess several areas within the stand that were representative of the population and they
also tried to assess the variations in the stand. That way you could improve the efficiency of field work and get enough good data despite the limited measurements.

To estimate the timber volume with the help of aerial photography, ie by measurement and interpretation of aerial photography, was tested in Sweden in the 1950s but had no practical significance until the late 1970s. The National Land Survey (then called “Lantmäteriverket” or LMV) began to use the technology to estimate the values of forest properties. The method is therefore often called "LMV" method. They used analog stereo instruments (figure 5.2) which could be used to measure the heights of trees, and they could also map the boundaries of stands with high accuracy. In the 1980s, as forestry companies used this method, it would improve its stand register. Several of the largest companies conducted new delineations (indelningar) of all or part of its forest assets by using the LMV method.

The LMV-method measures the stand’s average height and assesses the volume density and then calculated the volume. Usually, the calculation used Jonson formulas for full closed stands multiplied by the photo-interpreted stand density. In the beginning, however, a common relascope table was used with introversion plotted lines (Figure 5:37). Other functions for volume calculation have been used. Probably, however, the photointerpreter's ability to assess the crown closure is more important than which function to use.

In normal standwise inventory there is also a field inventory where it can be used to help quality control the photo-interpretation and supplement it with information which cannot be photo-interpreted, for example, timber quality. Most often each stand is visited but it is also conceivable that stands to be visited are selected using statistical methods, or those that are considered uncertain in their image interpretation results. The extent of the field control is dependent on the accuracy requirements. In volume estimate by LMV method there has usually been a mean error of stand level on about 15 - 20%.

If one is to estimate the timber volume over very large areas, an inventory/photo-interpretation can be instead done standwise using "point-lattice" method, ie to lay out plots systematically over the entire area. Each plot in the aerial images is then interpreted after which a control inventory can be done on a statistical sample of the plots in the field. The accuracy depends both on the representation of the error, and the standard error of the estimate of the single plot. The accuracy increases with increased plot number. By comparing the measured field and image estimates, potential systematic errors can be discovered and corrected. The Point Lattice method was used eg to estimate the total amount of timber damaged by the forest fire in Västmanland in 2014.
Figur 5.37. Relaskoptabell med ungefärliga massalutenheter.

Summary of Conventioneal- versus LMV method

Conventional method

- Usual personal computer or tablet used (stereoscopic used previously).
- Delineation on digital orthophoto
- Only stands delineated in the image (no estimates of the variables).
- At the boundary of digital orthophoto the result is a scalable real map database but respect to displacement of the treetops must be taken.
- A fieldwork map produced from the map database and used for field work. No interpreted data utilized.
- Image interpretation and field work is usually done by the same person.
- Short training of the photo interpreter.
- Inexpensive equipment.
- Simple process to get to work.
LMV-method

- Digital photogrammetric workstations are used.
- Digital stereo images are used. Delineation and interpretation of the stereo model
- Estimation of some variables by measurement and interpretation of images.
- Delineation and mapping done to scale
- A fieldwork map produced from the map database and used in the field work with the interpreted data.
- Image interpretation and field work is usually done by different persons.
- Relatively long training time required for image interpreter.
- Expensive equipment (software) required for image interpretation.
- Slightly more complicated process before the image interpretation can begin.

Litteratur


Lantmäteriet. 2013. Långsiktig Flygfotoplan. www.lantmateriet.se


9 Automated Digital Photogrammetry

Even aerial photographs can be used to produce three-dimensional information about the forest. Pictures taken in stereo can be used to create point clouds similar to those from laser scanning, with two important differences: i) points from aerial images also contain information on color, usually blue, green, red and near infrared, ii) because the camera does not "see through" the trees, there are virtually no points from the undergrowth and forested areas (Figure 23). In Sweden there is a good supply of aerial photographs thanks to Lantmäteriet’s photography program, and with the help of the new, national elevation model (NNH) which supplies the height of the ground elevation, which would otherwise be missing. This technology has not yet received the same circulation as laser scanning in forestry, but the research results are promising and of great interest.

Inspiration to the figures of Section 9.2-9.5 have been downloaded in Anders Boberg Introduction to photogrammetry [25], which is also an excellent source for those who want to know more about photogrammetry’s grounds.

Figure 23. 3D point clouds from laser data (left) and from the matching of digital aerial photos (right). The pictures show data from the same area. The profiles (bottom row), it is clear that the laser data provides more information on the ground and tree cover’s vertical structure. The digital photogrammetry, however, gives useful information on the tree cover height and also the color information in this case for example showing dead trees as blue dots. Photo: Jonas Bohlin, SLU. © National Land Survey, i2012/107. Reproduced with the author’s permission.
9.1 Color information from aerial imagery
Digital aerial cameras record usually the four colors blue, green, red and near infrared. Furthermore registered often panchromatic channel where light from a large part of the visible spectrum are used. The panchromatic band usually has a smaller pixel size and is used to improve the spatial resolution of the image. The colors blue, green and red can be "normal" color images produced where vegetation appears in green. One disadvantage of these images is that the blue band can be influenced by haze in the atmosphere, especially at altitudes above 3000 m. Often green, red and near infrared bands are used instead, but are displayed using the blue, green and red colors in the produced photographs. In these so-called “false color images” vegetation appears in red. An advantage of these images is that different types of vegetation can be identified more easily in the near infrared wavelength band.

9.2 Central Projection and orthophoto
In the type of digital camera that Lantmäteriet has used so far, as well as analog cameras, the image is a central projection. The image that this camera creates gives the landscape a so called “central projection”. Light rays from each point in the landscape pass through the camera lens (projection center) and continue to the detector, thus creating a mirror image (Figure 24).

Figure 24. Central Projection.

If the landscape is flat and horizontal, and the camera is looking perpendicular to the ground, the central projection (photo) will be a map-accurate picture of the landscape. In practice this ideal case never occurs, - camera tilt, down gradient and hills, hollows, trees and
buildings influence this. This has the consequence that the image geometry does not match the map's geometry. Figure 25 a) and b) show the scale of the image varies due to elevation differences in the terrain and tilting camera axis. Heights such as trees, houses and mountains seem to lean out from the center, known as radial displacement (Figure 25 c). The offset is proportional to the object height. Similarly displaced dips in elevation appear to sink towards the center.

There are several commercial software programs for digital production of orthophotos (map accurate images). The programs convert the images pixel by pixel. One can imagine the orthophoto as a grid of pixels which start with no content. The pixels in the original aerial photo should be given new positions in the orthophoto. This means that each pixel of the orthophoto will be filled with color from one or more pixels in the aerial photo. Orthophot pixels have known planar coordinates. The height of the corresponding point on the ground is obtained from a height model. If the camera position and angle are known, one can calculate the flight image for this part of the landscape that has been depicted. The color of that part of the flight picture is now transferred to the current pixel in the orthophoto. The conditions for making a digital orthophoto is that you know the orientation of the camera and have access to a digital elevation model, or create one from aerial photos.

9.3 Line Scanning and cylindrical projection
An alternative to the traditional camera is a line scanner. Instead of taking a sequence of two-dimensional images the landscape is scanned row by row. To obtain stereo images scanners are used that look in two, or more often, three directions: diagonally backward, straight down and slightly forward (Figure 26 a). Thus every point on the ground is photographed from three angles. Scanning lines are perpendicular to the direction of flight, but even in this case, differences in height and tilting camera axes will cause scale differences. As with the central
projection heights and hollows are displaced, but not radially from the image, but out to the sides perpendicular to the flight line (Figure 26 b). Such a projection is called cylindrical projection. The offset is proportional to the object height.

9.4 Stereo photogrammetry

The principle of stereo photogrammetry is the same as for human depth perception: two images from different angles, combined into a three-dimensional model with information about the distances to different objects. Commonly at least a 60% overlap between images is used, which by some margin provides stereo coverage across the image strip.

Figure 26. a) Line scan results b) cylindrical projection. In a cylindrical projection shifted tall items out to the sides, perpendicular to the flight line.

In the two-dimensional images the objects are displaced out from the center, see figure 25. The displacement is greater the closer the eye or camera object is located. An object is displaced in different ways in the two pictures, because they have different central points. The difference in shift is called parallax and allows stereo vision and range estimation. Aerial photos used for stereo interpretation overlap in the direction of flight. Orthophotos cannot be used because parallax caused by the topography has been removed. Nowadays, there are special computer screens that are used in combination with special glasses to enable stereo interpretation of digital aerial photographs. Digital photogrammetric workstations with such displays have replaced the earlier analog instruments to identify correct interpretation of aerial photographs.

Height and elevation can be calculated analytically for objects that are visible in both images of the pair. This is done by using the so-called parallax formulas. Figure 27 illustrates how the parallax formulas are derived when the camera axes are parallel to each other and perpendicular to the photographing base (distance between the two points when the camera made its exposure). It’s called
“photogrammetry’s normal case” but, despite its name, an ideal case in practice only occurs approximately. In the figure, the image plane is drawn in so-called positive position, i.e., showing the same side of the projection center of the imaged object (compare with Figure 24 where the image plane and the depicted landscape are drawn on either side of the projection center.

![Diagram](image)

**Figure 27.** An illustration of how parallax formulas are derived. The coordinate system xyz is the origin of the left image projection center O’. Object point P has the coordinates x, y, z coordinate systems marked ‘and” have the origin in each image center. The mapping of point P has the coordinates x’, y’, z’ in the left image, and x”, y”, z” in the right. Camera constant c is the image z-coordinate, and is negative. Photographing base is denoted b.

Similar shaped triangles gives

\[
x / x' = z / -c' \quad (Eq \ 7)
\]

which in turn gives

\[
x = x' * z / -c' \quad (Eq \ 8)
\]

p_x or x-parallax, is defined as x’- x”. Again, similar triangles give

\[
z / -c' = b / p_x. \quad (Eq \ 9)
\]
Combing Equation 9 and 8 finally gives

\[ x = x' \times \frac{b}{p_x} \quad (Eq \ 10) \]

In a similar manner one can show that

\[ y = y' \times \frac{b}{p_x} \quad (Eq \ 11) \]

and (because the y-parallax normally is zero and y' = y'')

\[ y = y'' \times \frac{b}{p_x} \quad (Eq \ 12) \]

and that

\[ z = -c \times \frac{b}{p_x} \quad (Eq \ 13) \]

The height z is the object's altitude. To determine the height above the ground the ground height must either be measured in the same way as above, or you need to have access to an ground elevation model.

9.5 Point clouds from stereo images

The conditions needed to generate point clouds from aerial photographs are i) the images overlap, ii) the camera interior orientation is known and iii) the camera position and angle are known with great accuracy. The camera's interior orientation is measured in the laboratory. The external orientation (position and angle) are recorded with GNSS and INS (section 3.4), but these values are often not accurate enough for a good image match to be made. Fine adjustment is done with the block triangulation. A common method for this is called “bundle adjustment”. Clearly visible and accurately measured points in the terrain, for example, painted white squares or manhole covers, are used to connect the images together into an image block. Bundles from these objects are used to fine-tune (adjust) the cameras external orientation.

When the conditions above are met, the images within the block are matched to the point cloud. There are some different methods for matching and the results are slightly different. Object-based matching requires that as many objects as possible are identified and matched. Pixel-wise matching (semi-global matching) is a newer method that ensures that every pixel is matched, resulting in very dense point clouds. After matching the objects, the x-, y- and z-coordinates are calculated using parallax formulas (section 9.4). The result is a point cloud very similar to that obtained with laser scanning, but one important difference is that the points can be complemented with color information from aerial photographs. Which principle that is used to do this varies between different software for image matching.
As the number of ground points is very low for the matching of aerial images, especially in forests, in practice it requires that you have access to an existing elevation model. Starting in 2009, Lantmäteriet has been conducting a national laser scanning in order to create a new national elevation model (NNH), and the possibilities of using stereo matched aerial photography is good.

9.6 Applications in forestry
Orthophotos are widely used as background maps in forestry, as well as delineation and updating of forest management plans. Even mapping in photogrammetric instruments has been used, particularly for forestry planning of major holdings. In the so-called "LMV" method, photogrammetric instruments were used to make the delineation of stands and to measure tree height, and to estimate the volume density (in Swedish "massaslutenhet") and tree species distribution. The height and volume density is then used to estimate the timber volume. With the introduction of digital images and digital photogrammetric instruments, this approach has become easier. It is not so widespread in the forestry companies but is mostly used by consulting firms.

There is a possibility to use orthophotos, for example to perform segmentation of stands, but different viewing angles in different parts of the image require special software. In addition, a two-dimensional image gives only limited information on forest height.

The technique of matching aerial images that is used to create 3D point clouds with the ability to depict the canopy similar to laser data provides new opportunities to automatically obtain forest information from aerial photographs. This is especially interesting for smaller properties because Lantmäteriet regularly photographs the country and distributes the digital images at a marginal cost.

If the points are converted into raster data, then they can be used for segmentation of populations with different heights and texture. It should also be possible to use 3D models of this kind in order to detect changes.

Point clouds from photogrammetry can also be used for estimates of height, timber volume and basal area in the same way as with the area-based method. A recently reported trial gave good estimation results when images from Lantmäteriets DMC camera with altitude 4800 m were used (Table 5).
Table 5. Accuracies at the stand level for the estimation of tree height and stem volume from point clouds created from the National Land Survey digital aerial imagery, from Bohlin et al 2012 [26].

<table>
<thead>
<tr>
<th>Variable</th>
<th>Metric used from point data</th>
<th>Accuracy at Stand level (RMSE)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Basal area</td>
<td>$h_{80}$</td>
<td>8.8 %</td>
</tr>
<tr>
<td>weighted mean height</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Stem volume</td>
<td>$h_{80}$ and a density metric</td>
<td>14.6 %</td>
</tr>
</tbody>
</table>

The experiments reported in Table 5 are very promising because it indicates that forest estimates can be made automatically with relatively inexpensive technology, with at least as good results as the manual methods. The condition is, however, just as for laser-based estimates, that there are reference plots of forest data that can be used as independent variables in the estimation. The forest used in the study above had tall forest arranged in well managed stands. Attempts of this kind must be repeated on more properties before firm conclusions can be drawn. It is clear that the point cloud from photogrammetry is a very valuable source of data for future forestry purposes.
6. LASER DATA FOR REMOTE SENSING

**LiDAR.** The technology to measure distances with laser is often called LiDAR from the English “Light Detection And Ranging”.

**Global Navigation Satellite Systems (GNSS).** GNSS is a generic term for all satellite-based navigation systems.

**Inertial Navigation System (INS).**

### Laser data in remote sensing

Laser scanning provides accurate three-dimensional measurements of vegetation and other surfaces. Airborne laser scanning has revolutionized forestry opportunities to efficiently produce forest maps with forest variables such as timber volume, basal area, tree height and trunk diameter. Through terrestrial (ground-based) laser scanning, it is possible to rapidly obtain field data with extensive information in comparison to traditional manual measurements. These instruments can be placed on a stative or can be a mobile system (e.g., on an ATV or a backpack with an operator). Information from terrestrial systems can be used to calibrate remote sensing data collected from airborne systems.

#### 6.1. Introduction to airborne laser scanning

Laser scanning has been established as a new and effective approach to forest data capture. Airborne laser scanning measures with decimeter accuracy the position of points both on the ground and in the canopy. With the help of special computer programs the ground height can be calculated. After this, statistical measures describing the distribution of the remaining laser points in the canopy are calculated. Using for example regression analysis the forest data from plots can be transferred to the raster cells which cover all forest land covered by the laser scanning. This so-called area-based approach was introduced in 2002 as a commercial approach to forest management planning in Norway and is now used by almost all forestry planning there. Foremost the area-based method provides measurements of tree size and number within each raster cell (stem volume, height, basal area, etc.). Delineation of stands and estimation of species distribution is done in Norway usually by manual interpretation of digital aerial photographs in stereo.
The Finnish state forestry centers have switched from a traditional, field-intensive forest mapping to making automated estimation of laser scanning. The area estimated by laser-based methods now amounts to 1-2 million hectares per year. Unlike in Norway, Finland tries to estimate tree species automatically, for example by combining the laser data with digital aerial photographs. The methods for automatically classifying detailed tree species from digital aerial photographs over large areas for operational projects, however, is still relatively uncertain.

In Sweden, most of the major forestry companies and some forest owners' associations tried to make laser estimates in the early 2000s, often on study areas of about 10 000 ha. The results were generally good in terms of variables related to tree size. According to a statement made by OL Forest Inventory AB the estimation accuracy (relative RMSE) of stand level with the area-based approach was 3-6% of basal medium height, 6-14% of the timber volume and 7-13% for basal stem diameter. These accuracies are much better than that achieved by traditional inventories made in connection with forestry planning. The accuracy of the estimate of the number of stems is about 12-24% (Brethvad and Iversen 2012).

The establishment of laser scanning as an operational method in Swedish forestry took momentum with the national laser scanning that the National Land Survey (Lantmäteriets) started in 2009. In 2011, the forestry company of Bergvik was the first major forest company in Sweden to order laser estimates for all its holdings based on Lantmäteriets laser scanner data. Since then, among others, Holmen has done the same and SCA has ordered the estimates for all of Norrland north of Sundsvall, even outside their own holdings. Even the National Property Agency (Fastighetsverket) and Sveaskog have ordered laser products from the consultancy that worked on Lantmäteriets laser scanner data.

In 2013 the National Board of Forestry was commissioned by the Government, together with SLU and other stakeholders, to make forest products over all of Sweden based on Lantmäteriets national laser scanning data. As a result, SLU made forest estimates based on raster data with 12.5 x 12.5 m size grid cells throughout Sweden except for the mountains. The variables created were estimated growing stock, tree biomass, basal area, basal area mean diameter and mean height. In addition, the Board of Forestry together with consulting companies developed additional products such as a 2 x 2 m tree height grid, and raster layers based on the DEM model from laser data showing hill shading, and ground wetness areas. These data layers can be accessed from the National Board of Forestry's website under the name of “skogliga grunddata” (in English translating loosely to “basic forest data”). The production of the first version of these forest data was completed in 2016, as all ALS acquisitions over the forest were not ready until then.
Initially the spread of the operational use of laser scanning occurred more quickly in Norway and Finland, especially for private properties. A contributing factor was probably that in these countries there was an established system of coordinating forestry planning for private properties over larger areas, while the state partially funded the plans.

6.1.1. Laser scanning’s development

Laser light is single color, directional light wave that is in phase. The first functioning laser was constructed in 1960. Today, lasers are used in several areas: in CD and DVD readers, printers, laser spectroscopy, the study of gases in the atmosphere, distance and speed measurement, topographic mapping, mapping of the sea depth in shallow areas and much more. Airborne laser scanning (ALS) uses the laser's ability to measure distance based on the time from when a laser pulse is sent out, until the laser light reflected from the ground or vegetation comes back to the sensor. The technology to measure distances with laser is often called LiDAR from the English Light Detection And Ranging.

The first attempt to measure tree heights with LiDAR was made in the former Soviet Union in the late 1970s. Measuring tree height profiles with airborne laser was also tested in Canada and the United States in the early 1980s (Nelson, Krabill, and MacLean 1984, MacLean and Krabill 1986). The first known studies of using an airborne scanning laser system for forest inventory in Sweden was carried out in collaboration between the then FOA (now the FOI, or the Swedish Defence Research Institute) and the Swedish University of Agricultural Sciences, 1991 (Nilsson 1996, 1994). These early experiments showed that stem volume and tree height could be measured by laser scanning. One problem with the experimental system used in this early test was the low accuracy in positioning. In the mid-1990s commercial laser scanning systems for airborne systems over land were developed. In these systems the GPS was integrated with inertial navigation (Inertial Navigation Systems, INS) which made it possible to determine the position of the laser measurement with 0.5 meter accuracy or better.

One of the first commercial systems was Swedish TopEye, developed by Saab in 1993. This helicopter-borne system produced data of high quality. The TopEye system was designed for detailed measurement of small areas and was widely used for the measurement of various types of infrastructure and to experiment with estimates of forest. The early attempts of helicopter-borne laser scanning did not lead to its use within operational forest inventory because the cost of data collection was perceived as too high. In Norway, Erik Næsset in 1995 attempted to measure forest with a scanning laser mounted in an airplane. The tests were successful, leading to the development of the so-called area-based approach for inventory at stand level with the support of field measured reference surfaces. In the late 1990s Finnish and Swedish research showed the ability to also detect and measure individual trees in laser data with high point density. An account
of the early development of the Nordic countries is given in the following reference\textsuperscript{1}.

Today, laser scanning is an established method for forest inventory. Examples of countries or regions where extensive laser scanning for forestry purposes are made are Norway, Finland, Austria, Spain, northern Italy, USA, Canada, Chile and Tasmania off Australia. Individual projects are in another wide range of countries, such as Nepal, Tanzania and Brazil to name a few. In 2011, Professor Erik Næsset from Norway received the Wallenberg Prize (also called the forest Nobel Prize) because he launched the area-based approach as an operational method for forest inventory. The year 2011 can also be counted as a breakthrough year for large-scale operational use of the method in Sweden, as the Bergviks project to underestimate its entire forest holdings with laser scanning started at that time. In Sweden, it is therefore 20 years from the first trials in 1991, to the full-scale deployment.

6.1.2. Basic characteristics of laser

This chapter describes the main principles of laser scanning and the main components used in today's systems. Foremost is the chapter on airborne laser scanning. A distance-measuring laser is placed in an airplane or a helicopter. The laser emits pulses to measure the distance to points on the ground or in the vegetation, and a scanning mechanism is used to distribute pulses in a wide swath beneath the airplane. For accurate positioning, the laser measurement points are required to be of high precision to register the distance to the point and the laser position and direction in which the pulses are transmitted.

6.1.2.1. Distance measurement

The method used for measurement of the distance (on the order of 100 meters or more) means that the laser emits a short (about 4-10 ns, corresponding 1.2-3 m), but intense light pulse. The pulse travels through the air, hits an object and is reflected by the object back to the instrument, which records how much time has passed since the pulse was sent out (Figure 6.1). The detector converts the light into an electric voltage where the signal strength is a function of time. Since the speed of light is known, the distance between the instrument and the reflecting object is calculated as

\[ S = \frac{v \times t}{2} \quad (Eq\ 1) \]

where \( S \) is the distance to the object, \( v \) is the speed of light and \( t \) is the time measured. The division by 2 is because light travels the same route twice: first from the instrument to the object and back. The method with the timing of the return pulse is common in both ground-based and airborne instruments.

\textsuperscript{1} Næsset et al. 2004
There is also a type of ground-based instruments that instead measures the phase shift of the returned signal from an amplitude modulated laser pulse. This method is very accurate but is only suitable for short distances.

6.1.2.2. Profiling laser

By placing a distance-measuring laser in an aircraft altitude profiles of ground and vegetation can be created. While the aircraft moves forward laser pulses are emitted with high frequency and hit the ground along a line in the plane's flight direction (Figure 6.2). The early experiments with laser measurement of forest in the 1980s were made with this technology and potentially it could be a cheap way to get forestry statistics for large areas.

6.1.2.3. Scanning

The methods for laser inventory of forest that are now used in practical forestry are based, unlike the profile measurement, on the scanning of the entire area to be inventoried. The height measuring laser is complemented by a scanning mechanism that distributes the measurements in a path under the aircraft (Figure 6.3). The scanning mechanism often consists of a
rotating or oscillating (wobbling) mirror. By flying several paths or strips which are adjacent to each other, it’s possible to cover larger areas.

Figur 6.3. Flygburen laserskanning.

Strip width is determined by the altitude and angle of the scan, that is, the maximum angle between the laser beam and the vertical. String width and air velocity then determines how large area can be covered per unit of time. The faster an area can be covered, the lower the cost, and for that would use a high altitude, air speed and scan angle. For a given combination of these parameters point density on the ground is determined by the pulse rate (number of transmitted pulses per second). Because users often want to have a sufficiently high point density and large area coverage per unit of time, development has moved towards scanners with a high pulse rate. A technical challenge is to combine the increased frequency of the greater power needed for scanning from a high altitude, because the laser needs more time between each pulse when the power per pulse is high.

The development, however, has been very rapid, and the pulse frequency of the current commercial systems are often several hundred thousand pulses emitted per second, compared with about 2000 pulses per second by the earliest commercial systems in the mid-1990s. A high frequency relative to the altitude results in a pulse does not have time to be received back at the sensor before the next is sent out, and therefore systems have also been developed that can keep track of multiple pulses while in the air.

For large scanning angles the proportion of the laser pulses that hits the vegetation instead of the ground increases. This means both lower coverage of the ground, and that part of the laser measurements used in forestry estimates become less representative. Often therefore there is a limit set on a useful scanning angle – which is approximately +/- 20 ° for laser scanning primarily for the production of ground models, and approximately +/- 15 ° for laser scanning primarily for forestry estimates.
6.1.2.4. **Positioning of laser point**

In order to calculate the position of the measurement point in horizontal and vertical direction the laser scanner’s position must be known as well as the orientation at the time of each transmitted pulse. The position is defined by three coordinates in space (x, y, z) and the orientation of the three angles called tilt, roll and yaw (Figure 6.4). Moreover, one must know the emitted laser pulse and direction relative to the instrument.

![Diagram](image)

**Figur 6.4.** Tipp, roll och gir (på engelska pitch, roll, yaw).

When the first experiments with airborne laser scanning were done, the technology was lacking to measure these parameters with sufficient precision, but in the mid-1990s, GNSS (Global Navigation Satellite Systems) started to be integrated with inertial navigation systems (INS). GNSS is a generic name for all satellite navigation systems, in the current situation the US GPS and Russian GLONASS systems. In practice, GPS is used more than any other system. GNSS provides good measurements of position and velocity, but the frequency is so low that multiple laser pulses can sent out between each measurement.

To measure the plane's orientation and fill in the gaps between the GNSS measurements used INS. An INS, or inertial system, often consists of three gyroscopes and three accelerometers. Gyroscopes measure the angular velocity around the three axes, and by integrating with respect to time, the change in orientation compared to the original orientation can be calculated. The accelerometers measure force and acceleration. Double integration with respect to time gives the position change from the starting point. GNSS and INS complement each other by the fact that INS fills the gaps between the relatively sparse GPS position and velocity measurements, while the GPS is used to correct for drift in the inertial system.

6.1.2.5. **Mätnoggrannhet**

For hard surfaces, the accuracy and resolution of the distance measurements primarily on tidtagningens accuracy and laser pulse length. Since the distance to an object is

\[ S = v \cdot \frac{t}{2}, \]  

(1)
where $t$ is the accuracy of the counting. The speed of light is approximately 300,000 km / s, so that a measurement accuracy of 1 dm requires a timing accuracy of 0.67 nanosecond ($1 \text{ ns} = 10^{-9} \text{ s}$). The speed of light is well appointed but slightly affected by air temperature, so a measurement accuracy better than anyone cm is difficult to achieve using airborne laser scanning, no matter how well time can be measured. The laser beam has low divergence, that is cohesive - compare the narrow beam of a laser pointer with the light beam from a flashlight. At large distances is the divergence nevertheless sufficient to pulse should see a small area on the ground rather than a point. A divergence of 0.5 mrad ($0.029^\circ$) and an altitude of 1000 meters gives a face (footprint) by approximately 50 cm in diameter. Larger divergence and altitude leads to the laser pulse, a larger hitting area, which affects the accuracy of the measurement negatively. Even large terrain slope and scanning angle can reduce measurement accuracy.

For hard surfaces, the accuracy and resolution of the distance measurements depend primarily on time-taking accuracy and the laser pulse length. Since the distance to an object is

$$S = v \times t / 2 \quad \text{(Eq 2)}$$

so, the accuracy of distance measurement can be approximated to:

$$\Delta S = V \times Dt / 2 \quad \text{(Eq. 3)}$$

where $Dt$ is the accuracy of the timing. The speed of light is about 300,000 km / s, which means that a measurement accuracy of 1 dm requires a timing accuracy of 0.67 nanoseconds ($1 \text{ ns} = 10^{-9} \text{ s}$). The speed of light is well defined but slightly affected by air temperature, so a measurement accuracy better than a few cm is difficult to achieve using airborne laser scanning, no matter how well the time can be measured.

The laser beam has low divergence, in otherwords it is coherent – as an example to demonstrate this, compare the narrow beam from a laser pointer with a beam from a flashlight. At large distances, the divergence is still sufficient so that the pulse should hit a small area on the ground rather than a point. A divergence of 0.5 mrad ($0.029^\circ$) and an altitude of 1000 meters results in measuring an area of about 50 cm diameter.

A part of the laser pulse can be reflected by an object that does not block the entire pulse beam path, while the rest of the pulse continues and is reflected towards items that will occur later in the path, for example, the ground. Many sensors can register more than one echo of the transmitted laser beam. For most sensors there must be at least one meter between echoes recorded from the same laser pulse.
6.1.2.6. Different types of scanners

The scanning mechanism usually consists of an oscillating or rotating mirror. A third type is a fiber scanner. An oscillating mirror oscillates between two positions and the laser pulses hits the ground in a zigzag pattern (Figure 6.5a). At the turn, the mirror slows down and the point density perpendicular to the direction of flight is higher in the strip’s outer edges.

One type of scanner rotating mirror is Palmerskannern (Figure 6.5b). The mirror surface is not perpendicular to the rotation axis, and the transmitted pulse direction changes with the position of the mirror. The pulses hit the ground in a circular pattern that moves forward in the direction of flight. The density is highest in the corridor's edges. A Polygon scanner (Figure 6.5 c) is also based on a rotating mirror. The mirror does not need to accelerate and brake, resulting in a smoother point density. Because of the aircraft's forward motion, the pattern is slightly angled towards the direction of flight.

The Fiber scanner (Figure 6.5 d) is made up of a bundle of optical fibers. A laser pulse is emitted into a fiber bundle in one end, where the fibers are assembled in a circle. At the other end of the bundle, where the pulse is emitted, the fibers are in a line. In this way the circular scanning motion is translated into a linear pattern. A rotating mirror divides the pulses between the fibers. The mirrors used are smaller than in other scanners and can therefore rotate faster. Point density is significantly higher in the flight direction than the strip width. One way to compensate for this is to allow the fiber-bundle emitting-end to swing back and forth so that the pulses from each fiber forms a zigzag pattern2.

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2 Petrie and Toth 2009a
6.1.3. Laser pulses interaction with ground and vegetation

On its way to the ground, the transmitted laser pulse can be reflected by the ground or objects above the ground such as tree crowns, trunks, other vegetation, rocks, buildings and more. A hard and compact surface results in a single distinct return pulse, but when the laser beam hits for example a tree crown or the edge of a roof, just a part of the pulse is reflected while the remainder continues travelling. Water has low reflectance and therefore has virtually no return.

Also a measure of the reflected radiation referred to as "intensity" is often recorded together with the coordinates for each detected laser return. Many laser systems use near-infrared light and a surface that reflects much light in this wavelength range (such as living vegetation) therefore appears "bright" in the intensity data. The intensity of the return pulses is difficult to interpret, because the amount of light reflected from vegetation depends on leaf angles, vegetation density, and other reasons.

Figure 6.6 illustrates how the reflection from different layers of vegetation and finally from the ground can give multiple returns from a transmitted laser pulse. The number of registered returns can vary between scanners. The forest often gives the first return from the canopy and the last returns...
from the ground. (There are also scanners that sample the whole laser return with high frequency, so called full-waveform laser).

![Laser pulse path to ground](image)

**Figur 6.6. Laser pulse path to ground.** This situation generates multiple returns - first from the tree, then from the bush and finally off the ground. The curve next to the image represents the return pulse’s intensity as a function of time, with peaks due to the different objects.

The height distribution of the laser points (the returns) occurs below the actual tree height distribution. This is partly because the laser beam penetrates the part of the tree crown before enough energy is reflected to trigger a measurement in the detector, and partly because the pulses not only hit the treetops, but also the tree crowns’ sides. Dense vegetation can also cause returns from the ground and vegetation to blend together so that the ground return will be partially masked. It may lead to the return being registered early and for the ground height to be overestimated.

The transmitted laser pulse duration is in the order of meters, but the uncertainty in the measurements against a hard surface should not exceed a few inches. Of great importance for the measurements is therefore the algorithm (calculation rule) that determines at what signal strength a detector will register a return. A common method is to register a return when the peak of the return signal has reached a certain percentage, for example 50%, of its maximum value (Fig 6.7). In this way timing is not affected by the pulse peak power, but only by its width. Peaks that do not reach above a certain threshold, the detection threshold, are not recorded at all.
A common method for recording the return signal is a peak in the signal which triggers an alert when it reaches a certain percentage of its maximum value, for example 50% (horizontal line in graph). Only peaks that reach above the detection threshold (dotted line) is recorded.

Laser systems are calibrated to give correct readings when hitting hard surfaces such as asphalt. The vegetation is however half permeable to the laser, and the fact that different systems use different principles for determining when a return pulse should be detected contributes to the values of vegetation of different laser systems being not directly comparable. The documentation available on the method used by the system to acquire and process the laser scanning data is often inadequate.

6.1.4. Processing of laser data

The laser returns recorded from the scan together form a so-called point cloud, which is a group of points which are coordinate-set in three dimensions (Figure 6.8 and 6.9). Most laser scanner systems also save the return signal intensity with the coordinates of the detected points. Before the data are used for further analysis they require some preprocessing. This is usually done by the data provider and is not specific to forestry applications.

Profile of laser point cloud where trees and buildings are clearly visible. The image is based on data from the National Land Survey. © Lantmäteriet, i2012/107.
Figur 6.9. Point clouds in three dimensions, seen obliquely from above. The points are colored from blue to red after rising height measurement. Through the area runs a stream. The image is based on data from the National Land Survey. © Lantmäteriet, i2012/107.

6.1.4.1. Stråkutjämning och inpassning mot kontrollpunkter

A number of factors that affect the positioning of the laser points, and an important source of error is the navigation system which consists of the GPS and INS. When an area is scanned it is usually in the strips which go back and forth, with some overlap between the lateral strips. They also tend to cross paths at the beginning, end and possibly the middle of the area (Figure 10). The uncertainty in navigation varies with time.

Figur 6.10. Outlays of the strips, cross strips, planar support (P) and the height support (H) in a scan area.

Data from different strips fit into a single seamless block via a process called strip equalization (Figure 6.11). Then the block is fit with the data
against the control points on the ground. Planar support is used to fit the data in the horizontal direction, and is composed of items that can be easily identified in the point cloud. Examples of suitable planar support are rooftops and ditches. Another possibility is to use white painted markings visible in the intensity data. Height support is used to fit the strips vertically, and should be done after planar alignment. Here open, even slightly inclined surfaces are desirable. Hard surfaces like asphalt and gravel are suitable because they give high accuracy in height measurements. ³.

Figur 6.11. *At strips equalization fit the different flight routes toward each other, whether in an average area or against a known surface.*

**Classification of ground points**

The erroneous data points removed are those that are high in the air as a result of reflection from such clouds, fog and birds, and those that clearly are below ground level. Low points can occur when a pulse is reflected in several steps so that the return signal is delayed, and the point seems to be farther away than it actually is. Classification of ground points can be made in different ways, two of which are described below.

One method was developed at KTH by Peter Axelsson and is used in TerraSolids software Terra Scan, and has received very widespread recognition [10]. To begin with, it uses a rectangular grid over the point cloud, with a cell size as determined by the user. In each cell, the lowest measuring point out is selected and classified as a ground point. The selected points are tied together in a network of triangles which provides an approximate representation of the surface. This data structure called a TIN (Triangulated Irregular Network) (Figure 12).

³ Olsson, Rost, and Reshetyuk 2013
Figur 6.12. *Ground elevation model in the form of TIN (dark gray lines). The lighter lines are contours. (After the picture of Robert Kropf at wikipedia. Licensed by the author under the GFDL).*

Now begins a new process of gradually adding new points in the network so as to densify and follow the surface details. Points in the cloud are examined one by one, and rejected or accepted as a new ground points according to certain criteria. One criterion is based on the distance between the point and the existing surface, another uses the difference in the slope of the ground surface of the new point added, and a third criterion is based on the maximum ground slope (Figure 6.13). What values are appropriate for these parameters is partly due to the topography.

Figur 6.13. *In Axelsson's method of land classification accepted a new point land point if the distance $d$ to the existing surface, the angle $v$ to the surface and ground slope $u$ below the selected thresholds.*

Another method is to first create a surface with the help of all points, where each point is given equal weight\(^4\). The surface falls somewhere between the ground and the canopy, and ground points are more likely under the surface than above. Each point is now given a weight which depends on its distance and direction from the surface (Figure 6.14). The lower the point is in relation to the surface, the higher the weight it gets, going down to a limit below which all points will get a maximum weight equal to 1. It also puts a distance above the surface where all points get a weight of 0 and therefore do not affect the surface. The weighting functions slope between the two extreme positions can be adjusted. The new weights

\(^4\) Kraus and Pfeifer 1998
are used to create a new surface, by points with higher weight "pulling" the surface to itself. The weights are updated again, and another new surface created. This continues a number of times - about 3-5 iterations may be required. After the last iteration the last computed surface determines which items should be classified as ground hits. This method is used in FUSION, a software for processing of laser data for forestry applications.

Figur 6.14. Weight p as a function of distance from the surface from Kraus and Pfeifers method\(^5\). Over the surface \((v>0)\) the weight decreases of above a certain value \(0\). Points below the surface \((v<0)\) get higher weight, the lower they are, but never more than 1.

6.1.4.2. Development of ground elevation model

Following the classification of ground points, the digital ground elevation model (or the Digital Elevation Model, DEM), is produced in the form of a TIN or raster. If the Axelsson method described in the previous section is used for ground classification, then there is already a complete TIN. If instead a method that only categorizes points into ground and other is used, for example Kraus and Pfeifer’s method describe above, it has a number of ground classified points that can be connected to a TIN. The benefits of a TIN is that all input ground points retain their coordinates in the plane, so that the accuracy of the point data is retained. However, it is a complex data structure that takes longer to create and process than a grid.

A grid can be generated from a TIN by is assigned a height to a raster cell from the TIN data structure has for the center of the grid cell. You can also create a grid directly from a collection of ground classified points either by using ground points within the grid cell or by an interpolation which also uses points just outside the grid cell. In a raster the points and their exact positions are not retained and hence this format does not taking advantage of the measurements’ full accuracy. Rasters are still the most common form of ground elevation model for use in forestry applications because they are very convenient to work with.

Laser data’s point density on the ground is very important for both the level of detail and accuracy of the final ground elevation model. Point density is

\(^5\) Kraus and Pfeifer 1998
affected by the pulse rate, flying altitude and vegetation. Dense vegetation means that fewer pulses hit the ground. The ground elevation model's accuracy is also affected by the accuracy of the distance measurements.

6.1.5. Estimation of variables for single trees

If the pulse density of the laser data is high enough then there will be many returns per tree crown. One can then detect the individual trees and estimate the characteristics for these. There is of course no limit to how many points are needed, but often at least 10 pulses/m² are needed, although lower densities can also work. Figure 6.15 shows the point cloud profiles from pine and spruce, scanned with about 50 pulses / m².

Figur 6.15. Point clouds profiles from pine and spruce, the pulse density of about 50 m². Image: Johan Holmgren, SLU.

The preprocessing of the data is done in the manner described in Section 6.1.4. Then the work can be roughly divided into the following steps:

• Assign height to all the laser returns over ground level

• Detecting individual trees in the laser data and delineate tree crowns using segmentation

• Calculate the measure from the laser data that describes the individual trees

• Connect the trees in the laser data with trees inventoried and coordinate set in the field

• Develop regression functions for variables to be estimated

• Applying functions on all detected trees.
Many of the processing steps above require special software. Analysis of individual trees in the laser data is therefore primarily done by researchers, as well as a few specialized companies.

6.1.5.1. Detection of trees

A common method for detecting trees is to first create a canopy height model (Digital Canopy Model, DCM), see Figure 6.16. Local maxima (peaks) in the crown elevation model is used as the basis of a segmentation. The goal is that each tree crown is represented by a segment (not to miss a few trees), and no crown should have more than one segment (to avoid "fake" trees). This means that the crown height model ideally should have exactly one local maximum per tree. A simply prepared crown height model that only follows the highest points in each grid cell does not usually meet this condition because it contains local height variations within the canopy. Therefore it uses some form of smoothing filter. The filter is adjusted to remove height variations within the canopy without removing the entire tree. The trees still missed are often small, hidden by the larger. Two trees standing close together can also be taken for one, while trees with broad crowns and several peaks can be interpreted as several trees.

![Figur 6.16. Crown Height Model with exposed tree tops. © Åsa Persson. FOA, Reproduced with the author's permission.](image)

A common method for the segmentation of individual trees is called “watershed segmentation” and is illustrated in Figure 6.17. In each grid cell over a certain height a "seed" is placed. The seeds then “climb” upwards in the direction of the steepest slope, until it reaches a point where all the surrounding cells have a lower value. This point is a local maximum and interpreted as a treetop. All seeds that climbs to the same point from the same segment, is assumed to correspond to the propagation of a crown. Alternatively, the segmentation is done in three dimensions, by some form
of clustering in the point cloud or by using so-called voxels, which is the term for three-dimensional pixels. There are a few studies which compare different methods of detecting individual trees.

Figur 6.17. Delineation of tree crowns using watershed segmentation. a) crown height Model filtered to give a maximum per tree. b) in each pixel of the crown height model is a "seed" that can climb the steepest path to a peak. A segment, or trees, defined as those pixels whose seeds climb to the top.

Alternatively, the segmentation can be made in three dimensions, through a kind of clustering of the point cloud or by using so-called voxels, which are three-dimensional pixels. There are a few studies comparing different methods of detecting individual trees\(^6\). The advantage of three-dimensional methods is that small trees in the canopy can be identified and that the segmentation of the canopy is more accurate even in height. The disadvantage is that the methods are often more complicated and that there is a greater risk that individual trees are divided into several segments.

### 6.1.5.2. Metrics of individual trees

Each segment may have an ID number, which is also given to the laser returns occurring within the segment. You can then connect all laser returns belonging to a tree and calculate different measurements of the tree. This is done for the trees. If segmentation is done in two dimensions, such as the above watershed segmentation, one must also find the crown lower limit in order to not get the points from the undergrowth. Among the benefits of laser estimation of individual trees it can be seen that species can be estimated from the crown shape, and that better information about the tree size distribution can be obtained. Some measures that have proven useful for estimating individual trees are:

- Tree height and crown area from segments
- Height Distribution dimensions in Table 6.1, calculated for individual trees, and the proportion of returns from tree crown

• Proportion of different return types, such as single-, first-and second returns from tree crown
• Texture Measurements derived from DCM and aerial photos
• Intensity (has no clear physical meaning if no calibration can be done, because it depends on many factors including system characteristics)
• Geometric measurement of the tree crown shape, such as the parameters of a parabolic surface that is adapted to a tree crown.

The measurements are used in later steps for estimating stem diameter, stem volume, tree height and tree species classification. Other measures that may be useful for tree species classification is color information from aerial photographs. (Holmgren, Persson, and Söderman 2008). One way is to project segments of the canopy on the aerial images using both color information and measurements from the laser data for each segment.

6.1.5.3. Linking field and remote sensing data

In order to estimate the tree variables from the segments, they must be connected to the field measured tree with known characteristics (height, diameter, etc.). The estimation with individual tree methods is not enough to know the sample plots position - one must also know the individual tree’s placement relative to the sample plot. It can be measured for example by ultrasound equipment connected to the autoclave (Haglöfs Postex) or compass. The position of the sample plot center should be measured with DGPS.

So that the pattern of laser measured tree and field measured tree should have the correct geometry, special programs are used to match field surfaces to the laser data on tree level. One method is to match two synthetic images with each other, and are summarized in the following points:

7 Olofsson, Lindberg, and Holmgren 2008

- Create a synthetic image of the tree pattern according to the laser data from the area around the sample plot
- Create a synthetic image of the tree pattern on the sample plot from field measurements
- Move and rotate "field image" to fit as good as possible with the "laser image"
- Connect the laser segment with field measured trees so that treetop distances are minimized.

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7 Olofsson, Lindberg, and Holmgren 2008
The synthetic images are composed of a dark background where the trees are represented by brighter spots. The brightness is determined in the laser image based on segment height, and in the field image by the trees measured diameter because you usually do not measure the height of the trees in the field.

6.1.5.4. Estimation of tree variables

Estimation of height, diameter, volume, etc. of individual trees can be done using, for example, regression or k-MSN, similar to the area-based estimation (Section 6.2). Regression models describing the relationship between the dependent (forest) and independent variables (from lasers or aerial photographs). The relationship is then applied to all detected trees within the scanned area.

Among the benefits of laser estimation of individual trees is that species can be estimated from the tree crown shape, and that better information about the tree size distribution can be obtained. To get unbiased results, meaning that the wood volume, number of stems, etc., on average, will be estimated correctly at the stand level, one must compensate for the trees that are not detected in the laser data. One method called semi-ITC achieves this by connecting all field measured trees to a segment, although in some cases it means that a segment has several field measured tree associated with it (Breidenbach et al. 2010).

6.2. Area-baserade metoder

6.2.1. Calculation unit

The area-based methods use a specific area unit upon which calculations are made, which is often grid cells in a grid covering the laser-scanned area. Grid cell size is determined typically based on a number of practical reasons, such as which field method was/will be used to collect reference data. The aim is often to estimate the totals and averages, for example, timber volume, basal area, average stem diameter and average tree height.

As reference data, circular plots having a size of a few hundred square meters are often used. The reason that a circular sample plot is used is that it is practical to measure in the field in all trees within a certain distance from the sample plot center. The circular sample surface area should be as large as the grid cell area that makes up the grid that covers the laser scanned area. The reason is that the variables should be calculated from laser data from the circular and square units which are as similar as possible.

What size is the best for raster cells depends on several things. They must be large enough to fit a group of trees in them and there should not be too many problems with edge effects, such as a tree outside the plot still has a canopy that covers part of the surface, which may result in the laser measurements indicating that there is a tree height, although there is a lack of trees in the field data plot.
If small plots are used, there may be errors in the positioning of the sample plot which would have a greater impact on the estimates. If large sample plots are used, the risk increases that the trees found in the sample plot / grid cell have completely different properties, such as the edge of a forest between the old and young forest. The size of the units (grid cells / plots) that is suitable also depends on the point density of the laser data.

The variables for each grid cell from laser data used in the models for estimating forest variables are calculated from the height distribution of the laser data reflections from the forest canopy and requires a certain number of observations so that calculations will be stable. It may be few laser reflections in the canopy if the laser data do not have a high point density, and if, at the same time, there is also a small number of trees within the calculation unit, and if these trees are small.

6.2.2. Variables calculated from laser data

To calculate the variables from the laser data for each unit, the height of the ground needs to be calculated first. The most common is that the height model is calculated as a grid with cells having a size determined by the distance between the laser measurement on the ground. Then the vertical distance to the ground is measured for each laser reflection. In the calculation of height measurements, one uses as a rule only vegetation hits above a set threshold, e.g., 2 m above the ground, to avoid returns from the ground and low vegetation which affect measurements. There may also be reason to calculate the measurements separately, for example, to calculate them separately for the first and last returns.

The term “height percentiles” refers to the height within an area, such as a grid cell, in which a certain percentage of points in the tree layer is found. More explicitly, 10% of the laser points are below the tenth percentile which is expressed as h10. Then 20% are below h20, and so on up to h100 corresponding to the highest point. The strongest correlation with forest height is usually around h90 (the 90th height percentile). The metric of h100 is often avoided because it is susceptible to occasional high trees and faulty laser points, and thus h100 is more dependent on point density than the lower height percentiles.

The proportion of all returned laser pulses reflected from trees is often highly correlated with crown closure. This percentage is called the vegetation ratio, here denoted $V$, and is usually calculated as the number of vegetation hits divided by the total number of laser hits. So called crown density metrics are calculated by dividing the height difference between the lowest and highest vegetation hits in a number equal fractions, often in intervals of 10. Then the proportion of returns over the respective fraction. Even the vegetation maximum height, average height, standard deviation and coefficient of variation are metrics that are sometimes used.
6.2.3. Factors that influence the laser returns distribution in the canopy

The technology used for the airborne laser scanning has been especially developed for the measurement of infrastructure and land surface. A flat surface is easy to define, and we get a clear return signal from it if we are sending a short laser pulse to the surface. The technology was developed for measuring ground surfaces even in tall vegetation by detecting multiple echoes from each sending of a laser pulse. It is important to remember that the technology was not developed to measure vegetation and instead treated the data from the vegetation as noise that would be filtered out. However, it was soon clear that airborne laser scanning could also be used for estimating forest variables because the laser beams penetrate the vegetation and describe the vegetation vertical structure.

The distribution of the laser reflection will depend on a variety of factors that are not always correlating with the forest variables that we want to estimate. We can describe the canopy of a variety of surfaces of varying size and location where each surface is not sufficient to reflect a laser pulse. If we have very frequent laser data (more than about five points per m²), we get a large number of measurements for each tree crown and it then becomes possible to geometrically model the individual trees. For area-based methods, we usually have sparser point density laser data and we model therefore not individual tree crowns geometry but use only the vertical distribution of the laser returns from the forest over the area of a grid cell.

We can think about how this vertical distribution arises. A laser pulse is sent down to the vegetation and a photo detector records the return of reflected laser light intensity. The system records the return pulse of an intensity peak which is detected above a certain threshold. Multiple returns can be detected from each sending of a pulse but the residual energy that is provided to detect the intensity peaks is then lower as the pulse has slowed in the upper layers of vegetation.

If an intensity peak is registered, it is due to a variety of system-specific factors: the algorithm used for detecting the intensity peaks, the sensitivity of the photodetector, the amount of energy in each sending of a laser pulse, and the wavelength used for the laser. Often, the system cannot register two return pulses which are close together from the transmitted laser pulse. This phenomenon also makes laser data from many systems less reliable for the measurement of shrub and field layers, as returns from these can interfere rebounds from the ground. For a given system, it is also dependent on the project: for example, a high flying altitude and a high pulse rate means a low energy of each laser pulse before it hits the vegetation.

If a high maximum scanning angle is used, the distribution of the laser reflections also depend on how far from nadir the pulse hits -- as a laser pulse that hits the vegetation at a very sideways angle must travel farther through the vegetation layers to reach the ground. There are also forest-dependent properties which have an influence: for example, distribution of
laser returns may be dependent on the tree canopy shape, tree crown density and crown height limit.

We can now imagine the situation that we underestimate average height and timber volume for raster cells with variables calculated from the vertical distribution of the laser returns in the vegetation. We note that the distribution of the laser returns depends on the distribution of reflective material in vegetation layers and a host of other factors.

When we estimate the average height of the trees, we usually estimate the basal area mean height. This weighted average has traditionally been used for forest inventory long before laser scanning was developed. It is calculated the same way as all weighted averages: Summarize the product of the measured values and the weight and dividing this sum by the sum of the weights. In this case, the weight of the cross sectional area of the tree (which we calculate from the value of the diameter with the assumption that the stem is circular). This average has been useful in forestry estimates because larger trees are important for eg timber volume.

There are studies showing that tree trunk cross-sectional area is proportional to the mass of the tree above the cross-sectional area. The probability that a laser pulse is returned from a certain tree is proportional to the tree crown mass which can be simplified as the size and density. So we have more laser returns from large trees than small trees. We would, according to this reasoning, expect a correlation between the basal area mean tree height and the average height of the laser returns in the canopy.

This relationship can be observed, but it complicates the situation that the laser light penetrates among the branches before a return is detected and the laser beams hits not only the tree tops but also the sides of the canopy. There may be a reason that a stronger correlation has been observed between the upper percentiles of laser returns for the height distribution and the basal area mean tree height.

It is important to note that a model is not used often to describes how the laser return distribution in the canopy is related to tree height distribution. If e.g. the basal area mean tree height is estimated in a two-layered forest, there is a high risk of underestimating because a large number of laser returns can come from the lower tree layer.

To estimate the timber volume, height measurements from laser data are often used, --- normally a percentile of height distribution of the laser returns in the canopy, along with a measure of density. As density measurements can the percentage of laser returns from the canopy be used. For estimates related to the density of the forest, we can use the connection between the total mass crown and the likelihood of a laser return from the canopy.
This relationship is also affected by several factors, of which perhaps the most obvious is illustrated by the difference between the proportion of the laser returns from the canopy of deciduous trees in winter and summer.

### 6.2.4. Planning of laser scanning

The variables we calculate from the laser data in the vegetation is a function of the vertical distribution of canopy and a host of other factors. Often we are interested in estimating forest variables describing tree stem properties eg, timber volume \( [\text{m}^3] \) per hectare. We examine empirically the variables calculated from the height distribution of the laser returns in the vegetation that may explain, for example, as much of the variation of the timber volume as possible and use regression or another method to calculate estimates.

If we do not get good estimates it can be useful to think about the physical characteristics of the laser measurement of vegetation and how the technology works. It is good to check as many influencing factors as possible. If we lack models to explain how different factors affect the estimates, it may be desirable to plan the laser data acquisition so that as many factors as possible are constant, such as using the same laser scanner systems, altitude, etc, for the entire project.

In Scandinavian coniferous forests dominated areas, laser data estimates have functioned at least as well for estimating forest variables from laser data acquired before leaf-out and after leaves have fallen off as compared to making estimates from laser data acquired during the summer. Laser Data collected from the period of ongoing leaf-out season or when leaves are falling off the trees should be avoided, because the amount of leaves may vary within the project area and between different species. In addition, laser scanning should be avoided when there is snow on the ground, as this affects the height model of the ground.

It is most difficult to control the factors that are determined by the forest that we measure. For this, there may be additional information from other data sources that can be used for stratification. If we have a stand register containing species information, it is possible to use different estimation functions for different kinds of forests, eg., forests dominated by different species.

### 6.2.5. Planning of field inventory

For both raster cells and plots, variables are calculated from the height distribution of the laser returns in the canopy. Laser and field data from test areas are then connected together so that it is possible to use the laser data to estimate forest variables in grid cells across the scanned area. The results can then be aggregated to the population level by calculating the average of all grid cells contained in a forest.

Sample plots for field data collection are laid out so that the forest variation of the area to be estimated is represented in the material field. A purely systematic sampling, for example in the form of a regular grid of sample
plots throughout the region, often results in too few plots in the stands of unusual character. One way to get a sufficient number of sample plots from all types of forest, without costs being too high, can instead be to first stratify the area, such as by tree species and/or age. Then there will be a certain number of plots in each stratum according to some objective method. Field and remote sensing (laser and aerial) data should be gathered close together in time. However, there are several advantages to delay the collection of field data until after the scan is made. Only then do you know what areas were really covered with useful data. Laser Data can also be an aid in the selection of field references.

If the sample plots field measured position is not consistent with the laser data the estimate quality can deteriorate significantly. Coordinates of the plots should be made with differential measurements with satellite positioning systems. At differential survey two receivers are used: one located on the sample plot and another that is placed on a well-measured point nearby. Information on the deviation between the measured and known coordinates at the reference station is sent to the device that is used on the sample plot so that the measured position of the sample plot can be corrected. For the attempts within laser scanning of forest that have been reported in the literature, the coordinate of plots has been set to 1 meter or better.\(^8\) Positioning accuracy is important when the forest is more heterogeneous and when small plots are used.

### 6.2.6. Estimation methods

Various estimation methods can be used, e.g. Regression\(^9\) and k-NN\(^10\). For regression estimates, usually only laser data are used and the estimates are usually not broken down by species. Instead, stands with different dominant tree species are estimated for different strata. Regression theoretically gives unbiased estimates, meaning that the value of the estimated variables average is correct. Since regression methods are model-based and allow interpolation (and within reasonable limits even extrapolation) they work with relatively little field data. The sample plots must be representative - for example, there should be a preponderance of plots where the forest is unusually dense in relation to their height. A limitation common when multiple regression is used is that each variable is then estimated separately, which means that unusual combinations of estimated variables may occur.

It has become common to use KNN methods where the laser data and aerial images are combined to produce species-specific estimates. The method is based on imputation, which means that for each grid cell is calculated based on forest data from sample plots that have similar characteristics as the laser data. This requires considerably more field data than regression, evenly spread over the entire range of variation. One advantage of the k-NN methods are that they measure multiple variables simultaneously,

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\(^8\) Næsset 2004; Packalen & Maltamo 2007  
\(^9\) Næsset 2002  
\(^10\) Packalen & Maltamo 2007
which means that there will be a more natural relationship between the estimated variable values, than with regression.

### 6.3. Future

The lasers that have been used commercially for airborne laser emits a laser pulse in a predetermined direction and a photo detector is used to detect the light reflected back from the messenger pulse. In this way the distance to the reflective object is measured. The emitted pulses are distributed over the area with the help of a mirror and aircraft movement. This means that the number of laser measurements per square meter depends on the laser pulse rate, altitude and scanners sweeping width. There are now commercial systems using an array of detectors instead of a single detector.

In order to understand how they are constructed, they can be compared with the detectors in a digital camera where there are several individual pixels. With these it is possible to split the returned signal in several parts from a pulse sent out. This makes it possible, from a given altitude, to gain much higher resolution laser data compared to using conventional laser systems, which means higher resolution for the same cost as before.

For forestry applications, it is important to quite often be able to collect 3D data for large areas, which has been a limitation with airborne laser scanning using linear detectors. If we use multiple detectors in a matrix, laser data with higher resolution is collected within the same amount of time as before, for example, if an array of $32 \times 128$ photodetectors are used, we can get 4,096 measurements from each laser pulse sent out.

We can, as previously done, use a scanning system, but each laser beam is divided into sub-beams, with each pulse return detected by a single detector in an array of detectors. The laser light is split, the energy that comes back is weaker for each detector. It is possible to use this technique because the detectors are very sensitive - so sensitive that they can detect single photons. That so sensitive detectors are available also means higher flying altitudes can be used because so little energy needs to come back to the sensor. The amount of light energy that comes back is inversely proportional to the square of the distance. There are so-called photon-counting systems that can be used from approximately 10 000 m flying altitude.

There are two different types of photon-counting systems that can be used to efficiently scan large areas of land. They have various types of detectors. A sort of system has been photo-multiplier tubes (PMT) which are very sensitive photodetectors. However, it is difficult to manufacture these for detecting near-infrared light which is the standard for the measurement of vegetation. Instead green lasers are often used which unfortunately do not reflect equally well from vegetation.

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11 Stoker et al., 2016
It is possible to measure time very accurately when the photons arrive at the detector and the sensor can detect multiple photons that are close together in time, which means that multiple returns can be detected for each detector and sent laser pulse.

Another kind of system has Geiger-mode Avalanche Photo Diodes (APD) for detection. Geiger-Mode is a way to use the APD so that a fast electrical pulse of several volts can occur when a single photon comes to the sensor. It is not as difficult to manufacture such detectors for near-infrared light. They can also measure time very accurately when a photon comes to the sensor but the sensor cannot usually measure the photons that are too close together in time, for example, if a strong signal coming from the upper part of the crown of the tree, the lower part of the tree cannot be detected from the same pulse. 12. Both types of detectors are extremely sensitive and, therefore, noise such as caused by sunlight arise, but because so many photons are detected, there are algorithms that can effectively filter the data.

There are now demonstrations where the new systems have shown an efficiency (per unit area) that are up to 30 times compared with traditional technology for airborne laser scanning. 13.

6.4. Terrestrial Laser Scanning

6.4.1. Terrestrial Laser Scanning and other ground based methods

Ground-based laser scanning and terrestrial laser scanning (TLS) is a technique widely used in the industry today. In the past decade there has been an increased interest in using these methods in forestry. There are mainly two aims: to get a quick field inventory and to get more information from the field survey. Considering the weight and measurement speed of the instruments today TLS solutions are nearly as fast as manual measurements. With technological development systems can probably be built that are both easy to carry and faster than manual measurements. In addition, these systems can be mounted on vehicles and thus cover much larger areas than usual. Because TLS measurements sample from the entire tree including trunk, branches and needles / leaves models can be built with high accuracy, on everything from biomass and basal area to the stem volume and stem shapes.

Although the research is new, there has been technological development regarding extraction of forest information from TLS data. For example, Tanaka et al (1998) used a system in which a laser plane was projected on to the forest area and the 3D structure was obtained by triangulation. The parameters extracted were stem position and diameter, among others.

A widely used technique of TLS systems is to produce data in point form as a list of coordinates and intensity. Hopkinson et al (2004), demonstrated the

12 Stoker et al., 2016
possibility of using such systems in forest calculations to obtain variables, such as height, volume, and diameter at breast height (DBH). The study used manual methods to extract information but indicated how these forest variables could be obtained automatically in the future.

Other examples are: Tanaka et al (2004) and Hosoi and Omasa (2006), showed that methods of extracting leaf-area-index (LAI) from the TLS data; Gorte and Winterhalder, (2004), extracted branch structure of individual trees and Pfeiffer and Winterhalder (2004), made a more detailed measurements of stem form and measured characteristics such as ovality, and Thies et al (2004) implemented methods to extract taper, curvature and leaning from TLS data.

6.4.1.1. Equipment

A TLS instrument weighs today less than 20 kg and is equipped with a tripod, see Figure 6.18. Some instruments are mounted on vehicles and the technology is called Mobile Laser Scanning (MLS). The wavelengths range from green to infrared. The sampling rate is around 1 000 000 points / second today and technology is developing continuously. TLS instruments can be divided into a number of categories: short, medium and long distances; panoramic scanning and camera scanning; phase measuring system and pulse measuring systems.

Systems built for short distances work within the radius of 50-100 m while the systems for long distances can reach more than 500 m of highly reflective surfaces. Panorama types of instruments usually use rotating mirrors to cover a hemisphere with different sample patterns while the imaging system takes pictures like a conventional camera with the addition that the distance to the object is saved along with the intensity of each camera pixel.

Phase measuring systems transmit a continuous wave (CW) which is amplitude-modulated. The phase difference between the reflected wave and the transmitted wave measured. To determine the ambiguity problem that the distance to an object can be longer than the wavelength used, a number of frequencies and the longest wavelength are factors used to determine the maximum possible measurement distance.

Pulsmätande system eller "time of flight" (TOF) system mäter tiden det tar för en laserpuls att färdas från instrumentet till föremålet och tillbaka. Eftersom ljusets hastighet är känd går det att extrahera avståndet. Se Shan och Toth (2009) för mer information om lidartekniker.

Pulse measuring systems or "time of flight" (TOF) systems measure the time it takes for a laser pulse to travel from the instrument to the object and back. Since the speed of light is known it is possible to calculate the distance. See Shan and Toth (2009) for more information about LiDAR techniques.
Figur 6.18 The Leica terrestrial laser scanner can be used to scan buildings, rock formations, forests, etc., to produce a 3D model consisting of a list of coordinates. The instrument can direct the laser beam in a variety of angles: the head rotates horizontally and the mirror rotates vertically. The laser beam is used to measure the distance to the first object in the beam path. Source: Wikimedia Commons.

For imaging at close range, flash laser can also be of interest in the future. Such a system sends out a very short pulse that illuminates the entire surrounding area, unlike a conventional laser scanner that sends out a pulse in a certain direction with a small divergence (spread). The time for the laser pulse which is sent out is measured with a matrix of photodetectors and the time is also measured for the light which is reflected back. In this manner, a 3D image is created without a scanner needing to split single laser pulses over the area to be measured.

Therefore small and inexpensive systems can be constructed and these are sometimes called "time-of-flight cameras" because they can be compared with cameras which measures the time it takes for light to travel to the object and back to the sensor. One advantage is that the whole scene is measured simultaneously as opposed to a scanner where it takes some time to scan the surrounding objects. It can be an advantage if you use mobile systems and / or follow moving objects. Additionally, the entire scene is scanned with a high frequency.

One problem is that much energy or highly sensitive detectors are needed to measure objects at long distances because the energy sent out is distributed over a large area and not that of the scanned system only a small area ("footprint"). We can expect that in the future there are small systems that can be hand held or that we place on eg drones or forestry machinery.

6.4.2. Data characteristics
The output from the TLS systems is usually in the form of a 3D point cloud where each position also has an intensity value (sometimes color values red, green and blue). Some systems also save other metadata such as
vegetation class and returns number of the pulse, etc. Data can be saved in text format or in a binary formats such as LAS. (read more on the LAS format on http://www.asprs.org/). The text-based formats are usually lined up in columns with a coordinate or color per column. For small amounts of data it can be read into text based files into a common spreadsheet software but often the data sets are so large that other analysis tools are required.

The TLS system measures the distances to surrounding objects with millimeter resolution based on the emission of laser light and detection of the reflected signals. Depending on the measurement resolution there can be several points per square centimeter covering an object which is ten meters from the sensor, which gives a very detailed map of the trunk and branches of a tree, Figure 6.19.

Figur 6.19   Examples of a TLS test area. The round hole is where the instrument has been, so that the self-shading has occurred. Behind the trees there will be shadows where the laser beams do not reach.

When forest information are to be extracted from TLS data, statistical and mathematical software are necessary that can read LAS file format, or the text-based formats. Alternatively, a self-made program developed that reads and analyzes the TLS data can be created and used. A variety of visualization software may display the TLS data. An example is available online is fugroviewer.com.

6.4.3. Using TLS for field inventory

TLS can be used for field inventory in the same way as the traditional manual methods. The instrument can be placed in the middle of a sample surface and any size sample plot can be selected using the distance data from the scan. Both systematic and random sampling designs can be used. It is also possible to use line-intersect sampling or belt-transect sampling – in that case the scanner could be located on a vehicle or a portable frame.

Om TLS-instrumentet används vid provytesampling finns det två vanliga metoder: ”single scan” och ”multi scan”, figur 6.20. I ”single scan”
If a TLS instrument is used in plot inventory, there are two common methods: "single scan" and "multi-scan", Figure 6.20. In "single scan" method the instrument is placed in the middle of the plot and scans only the side of the tree facing the sensor. The "multi-scan" method places the instrument in a variety of positions in the plot and therefore provides many views for each individual tree. Laser points from the different views must be rectified to a common coordinate system before use.

Figur 6.20 (A) Examples of "single scan" method. (B) Example of "multi-scan" method. Position for TLS instrument is marked in blue and the sample plot is marked by a circle. Trees positions are marked with green.

The advantage of the "single scan" method is the measurement speed and ease of use. The instrument needs only one position per sample plot and there is no need for rectifying the second sensor positions. The downside is that some trees will be shaded by other trees, so then some trees will be missing in the analysis. There is a statistical method which can take into account the missing trees.

The advantage of the "multi-scan" method is that point density becomes higher, allowing a higher degree of detail. It is possible to cover all the trees if the instrument is placed in many different positions. The downside is that reflectors are needed for rectification among all the points from all the instruments positions, and that the measurement takes longer.
6.4.4. Automatic extraction of forest variables from TLS data

Output from the TLS instrument contains only the positions and intensities / colors in 3D space and not variables such as basal area and stem volume. To calculate (or extract) forest variables from TLS data, signal and image analysis techniques are used. The most common technique is to extract the tree geometry from the 3D point cloud. If for example a cylinder is adapted to the data points that are far down on the trunk of a tree, basal area and DBH can be calculated with a given accuracy dependent on the quality of the algorithm. If multiple cylinders are adapted along the trunk of the tree, a stem profile can be calculated and if the branches are modeled, the number of branches on the trunk can be calculated. The amount of laser point hits in the branches and foliage can be used to estimate biomass.

Different algorithms are differently refined from simple models of DBH to advanced models where virtually every twig is represented. However, there are four sub-steps that usually are represented in most of the algorithms:

- Calculate a digital elevation model (DEM)
- Separate data points for each tree in the examined the sample plot
- Classify data points for each tree in categories such as trunk, branches and leaves / needles
- Model the detected trees using points for the different classes.

6.4.4.1. Calculating the DEM

Some of the algorithms for tree detection and modeling of TLS data needs a digital elevation model (DEM) as input data to estimate the DBH of the trees in the test area. Many of them create a raster where the height values of the minimum data point within each cell is recorded as land elevation at that position. Simonse et al. (2003), for example, created a grid of 50 × 50 cm 2 cell sizes. Many algorithms have been created to extract them from the airborne laser scanning; some of them can be adapted for TLS data. One of the earliest was developed by Axelsson (2000) and is based on a triangular irregular network adapted to the paragraphs below. The process is iterative and refined step by step where points are rejected if they give an unreasonable ground model.

Isolate data points for every individual tree

There are a number of techniques to separate each individual tree in the TDS data from the test surfaces. One method is to detect the part of the stem which is closest to the ground to find an initial position. It is possible, for example, cut out a section of the point cloud located at chest height Figure 6.21.
Figur 6.21  A cross-section at the height of 1-1.5 m from the ground level for a TLS data sets. The data points are projected into an image. A large number of points in each pixel provides a high intensity. Stems in the images can be seen as circles surrounded by branches.

From these cross-sectional images it is possible to find positions for tree trunks. Some methods are very simple measuring only point density in different parts of the image. In those places where there are many measuring points algorithms assume that there is a tree in the middle. Other algorithms are more advanced and are looking for circular shapes in the image. Trees positions assumed to be in the middle of these circles. Hough is such an image-processing algorithm that is looking for geometric shapes in images such as circles.

Once the trees' positions are known, the data points closest to a strain believed to belong to the tree. All data points that are below a certain level can be cut away so that the data points from ground level are not included in the analysis. For the algorithms that have been detected circles tribal forms can be cut dataset belonging to tribes where most of the branches are filtered out. For these cuttings it is possible to model the stems more closely.

One way to classify and demarcate tribes, twigs and pine needles / leaves is to find the characteristics of the spatial distribution of the small parts of the point cloud. They can be classified as spherical, flat and linear using the eigenvalues, Figure 6.22. Flat areas of TLS data can be assumed to be parts of a tree trunk or branch, while coarse linear parts is probably small twigs. By linking many flat parts to the surfaces it is possible to find points belonging to tree trunks, Figure 6.23.

Figur 6.22. Bild av en del av ett träd där de TLS data delarna som tillhör stammen är markerade i grön. De data delarna som ligger i ett plan område anses vara del av stammen.

Figur 6.23. Picture of a tree where the points from a TLS data sets stored in cells (volume pixels or voxels) with a high point density saved as a higher intensity. From the data it is possible to find trunks and thick branches.

6.4.4.2. Classification of data points for every tree in different categories

Once the data points for each tree are defined, they can be classified into trunks, branches and foliage. For both the voxel based and surface-based methods it is possible to find the first trunk lying at the bottom, and then follow the stem upward and connect the ramifications that arise. The line starts at the ground classified as trunk while the others become branches. The methods used are within the field of mathematical morphology where Dijkstra's "minimum spanning tree" is a common technique, Figure 6.24.

Figur 6.24 If the data points from a TLS measurements are saved in a 3D grid (a voxel volume) as in the picture to the left, it is possible to look for the trunk and branches by connecting the different volumes to a so-called "minimum spanning tree" as in image right.
6.4.4.3. Model the detected trees

When forest variables to be extracted from the point cloud, it is usually an advantage if the geometry of the scanned trees modeled figure 6.x9. Different studies have different refinement of these models: from the simple measurement of DBH to models with twigs and leaves. The most common technique is to adapt circles or cylinders to the data points that belong to the trunks and branches by nonlinear or linearized least squares. Some studies use robust adaptation of the cylinders, for example, Liang et al. (2009.2012) who used "Tukey's estimator".

There are also a number of other techniques to model the tribes. For example, it is possible to find the edges of the trunk of an image. By the distance to the strain is known it is possible to get an estimated diameter.

Some techniques model tree trunks with triangular irregular network or various kinds of splines. These more advanced models that show every bump on the surface can be used to obtain a volume of trunk alternatively to obtain an equivalent trunk diameter giving the same volume as the crooked model. For these advanced stem models it is possible to get the measure of quality, such as straightness, taper and ovality.

Figur 6.25. Examples of the stem profile of a tree that is modeled as circles extracted from TLS data.

6.4.4.4. Cirkel/cylinder adaptation of stems by regression

In the regression of circles to a point dataset minimizes the residuals of the points to the circumference of the circle. The position and size of the circle that gives the least error is selected as an estimate of the tribe, Figure 6:26. One problem with the regression is the sensitivity of the data points that unilaterally located on the outside of the circle. If stock items are not properly filtered in advance without a lot of branches and twigs are represented in the selected dataset errors in position and size to be included in the estimates of the stems. There is a need for a robust circle adjustment method for stems to be estimated with a good accuracy.
Figur 6.26. Examples of a circle adapted to a synthetic dataset TLS a strain with branches. The branches affects the position and size of the circle, thereby giving the errors affecting the estimation of the tree trunk. There is a need for a robust circle adjustment method or a good method to filter out branches before the stem's estimation.

THE FOLLOWING NEXT FEW PAGES ARE NOT TRANSLATED TO ENGLISH AND ARE NOT REQUIRED READING. SKIP AHEAD TO THE MOBILE LASER SCANNING SECTION.

6.4.4.5. Cirkelanpassning av stammar genom RANSAC algoritmen

RANSAC är en förkortning för ”Random Sample Consensus” och är en iterativ algoritm. Den är baserad på det faktum att det existerar datapunkter som tillhör modellen (inliers) och de som ej tillhör modellen (outliers). Om ett antal slumpmässiga punkter i ett dataset väljs så finns det en viss sannolikhet att alla tillhör modellen. Ekvation 6.x1 ger det antal iterationer som krävs för att få åtminstone ett sådant fall.

\[ N = \frac{\log(1-p)}{\log(1-wn)} \]  

där \( N \) är antalet iterationer där åtminstone en bra modell är hittad med sannolikheten \( p \). Antalet valda datapunkter är \( n \) och \( w \) är sannolikheten att en punkt tillhör modellen. Parameter \( w \) ges av ekvation 6.x2.

\[ w = \frac{M}{D} \]  

där \( M \) är antalet ”inliers” i modellen och \( D \) är antalet punkter i datasetet. Vanligtvis är denna kvot ej känd utan måste estimeras från ett dataset. Genom att studera ett typiskt dataset är det möjligt att erhålla en grov uppskattning av denna kvot.

Skärningen mellan två sådana linjer ger centrum för cirkeln om två korda används.
För att estimera parameter \( w \) i ett TLS dataset måste kvoten mellan antalet punkter i stammen och totala antalet punkter inkluderat grenarna bestämmas. Antal iterationer bestäms genom att sätta parameter \( p \) i ekvation x1 till en hög sannolikhet och genom att multiplicera parameter \( N \) med en säkerhetsfaktor beroende på hur många bra modeller som behöver hittas för att erhålla ett bra estimat.


Figur 6.x12. I varje iteration i RANSAC algoritmen hittas ”inliers” med en given tolerans för den valda cirkeln. Om antalet ”inliers” är större än ett given tröskelvärde behålls modellen för vidare beräkningar.
Figure 6.x13 Gör en cirkelanpassning för varje modell med ett högt antal ”inliers” i RANSAC algoritmen, genom att använda de utvalda punkterna exempelvis genom regression. Se om felen är mindre än i föregående iteration. Behåll modellen med de minsta felen.

6.4.4.6. Cirkelanpassning av stammar genom Hough-transformen

6.5. **Mobile laser scanning**

A mobile laser scanner system consists of the following components:

1. laser scanner,
2. positioning, and
3. other equipment such as computer and batteries.

All parts including a scanner, for example, mounted on a frame that is attached to a sling and scanner mounted on top of the frame with a clear view in all directions (see Figure 6.30). The mobile system resembles an airborne system so that a positioning needed to continuously measure the position of the sensor. For an airborne system, a satellite navigation system combined with inertial (accelerometers and gyroscopes). Using inertial changes in movement and orientation calculated with high precision and high frequency system, but affected by the operation, which means that the wrong absolute position calculated after only a few seconds. Therefore, data from a satellite navigation system (which does not have as high frequency but a position calculation without operation) to continuously calibrate the absolute position.

Figur 6.30. An experimental mobile laser scanning system with laser scanner, stereo camera, inertial navigation and computer for data logging and wireless communication network to other devices.

For mobile systems that are portable or mounted on forestry for the problem of signals from satellites often are not as good under the canopy. There are various ways that this problem can be solved. One way is to use
Stereo cameras that collect a sequence of images with high frequency. Algorithms can be used to automatically identify the pixels depicting the object in subsequent images and 3D coordinates can be calculated for points on these items. By minimizing the distances between points that are associated to the same object is calculated camera position at a time relative position of the camera at a different time. In this way, the sensor movement is expected in 6 dimensions (x, y, z and three rotational angles). This calculation can be used as the start value for further calibration.

From the system, we also get data from a scanner that sweeps 360 degrees several times in every second and sends out several hundred thousand laser pulses every second. If we have two point clouds from the subsequent sweep, the algorithm "Iterative Closest Point" (ICP) used. This algorithm consists of the following steps:

1. Find correspondence - that associate points in pairs to each other,
2. Find a function that minimizes the distances between the points of the two data sets,
3. Calculate new coordinates of the feature from the previous step,
4. Repeat steps 1-3 until the error is small, or the maximum number of iterations.

The easiest way to associate points to each other is to connect the points that are closest to each other. There are other ways to associate points e.g. "Point-to-plane" in which a surface is determined by the surrounding dots. The distance between the pairwise registrations is then calculated from a point in one point cloud, and along the normal to the surface of the second point cloud. In this way the plane surfaces slide over each other so that the structures are better matched.

The algorithms used to detect and estimate the tree trunk diameter similar to the algorithms used for stationary ground-based laser scanning (TLS). However, it is not certain that it is best to use the exact same algorithms. One advantage of the mobile laser scanning is that the trees are measured from many different directions, which means that the risk is reduced for zones that are obscured by other trees and there is a greater chance of trees upper parts, such as tree top, can be measured well from any direction.

The errors that come from calculation of the position of the sensor means that the point cloud’s accuracy will be lower, especially if satellite navigation systems are used to support the INS in the dense forest. Estimation Errors have proved to be greater than what is common in
statistical measurement.  

If many measurements on the tree trunk and the error is normally distributed, it is not certain that the estimation error becomes large. A comparative study showed that data from a handheld laser scanner was better compared with data from static laser scanning to estimate trunk diameter, a probable cause is that the measurements from the handheld system were around the entire trunk. 

As examples of how data from mobile systems can be analyzed is the "region growing" algorithm. There is an algorithm to segment the point cloud so that each 3D point given an ID number is defined as belonging to a particular object. First a threshold is determined for the mean square deviation (s) from a surface, and then the maximum angle difference (α) between two normals. The two variables \( r \) and \( α \) are calculated for each 3D point using a local area. This neighborhood may be determined in two ways: K-Nearest Neighbors (kNN), and Fixed Distance Neighbors (FDN). For kNN the k nearest neighbors to the current point are selected.

If we assume that the point density indicates the measurement accuracy, which is often the case because the point density is inversely proportional to the distance between the scanner, this is a good way to define the calculation area normals because a larger area selected for the longer distance of the sensor. It is also possible to avoid cases where the algorithm collapses because of the low point density in a certain area. For FDN selected all neighbors within a certain distance. This method is suitable for point density is constant within the measuring range.

The following steps are used:

- (1) A surface is calculated using points and the normal and the mean square deviation of the surface is calculated,

- (2) The point that has a surrounding surface with a minimum mean square error is selected as the first starting point,

- (3) The items found in the area included in the region of the angular difference between their normals and the current punkterns normally not greater than \( α \). The points have a mean square error which is less than \( r \) is placed in a list for possible start values,

- (4) If the list of possible starting values are not even going to the next available start value and repeat steps 3,

- (5) Include the region segmentation result and go to step 2.

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14 Liang et al. 2014
15 Bauwens et al. 2016
16 Rabbani et al. 2006
This is repeated until the list of items not included in the segmentation is empty then the list of 3D points that have been segmented are sorted in order of magnitude. One advantage of this algorithm is that it has so many parameters and these parameters can be explained physically. Depending on how the parameters are selected there may be an over-segmentation (many segments) or under-segmentation (few segments). In steps after segmentation geometric objects are modeled, and it may be easier to detect under-segmentation compared to the over-segmentation\textsuperscript{17}.

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7. RADAR REMOTE SENSING OF FOREST

Radargrammetry. Can be deduced from the optical term stereogrammetry but applied to radar images. Its fundamental principle is the stereoscopic viewing of an object from somewhat different angles.

Polarization. Is a parameter applying to waves that specifies the geometrical orientation of the oscillation, which in radar remote sensing is generally either Vertical or Horizontal.

Synthetic Aperture Radar (SAR). The Radar technique of synthesizing a large antenna by moving a smaller antenna along a path, in order to retrieve a 2D radar image with high resolution in both azimuth and range direction.

SAR interferometry. A remote sensing technique that enables accurate measurements of geophysical parameters, which mainly are related to height changes.

7.1. Radar basics

Each remote sensing technique has its pros and cons, but most ones are influenced by the prevailing weather conditions. Radar is a technique that is barely influenced of cloud coverage or rain. Radar (acronym for RAdio Detection And Ranging) is an active sensor for detecting, locating, tracking, and identifying objects even at a considerable distance. Generally, three basic types of radiation are commonly used in remote sensing. 1) Radiation emitted by the object itself, due to its material properties and physical conditions, for example thermal radiation. 2) Diffuse scattering of natural illumination from an incoherent radiation source, for example the sun. 3) Backscattered radiation from artificial coherent sources, for example, radar and laser systems.

The radar system is a sensor transmitting and receiving electromagnetic energy at micro wave frequencies, which means wavelengths much longer than visible light or thermal heat. The radar signal is capable of sensing the vegetation or other features on the ground regardless of sunlight and cloud cover. This is a great advantage compared with passive optical sensors, which depend on reflected solar radiation and relatively clear skies.

Radar sensors are partly characterized by their operating frequencies. The fundamental relationship between wavelength (λ) and frequency (f) in free space is determined by:

\[ c = \lambda \times f \]

where \( c \) is the speed of wave propagation, which is the speed of light \( (3\times10^8 \text{ m/s}) \). Thus, knowing the wavelength, the frequency can easily be...
calculated and vice versa. The wavelength from a radar system determines the extent to which the signal is attenuated by an object, e.g., the vegetation. In general the attenuation increases as the wavelength decreases. For example the effects of rain and clouds are negligible using a wavelength greater than about 3 cm.

A radar sensor can be mounted on almost any type of platform, and radar systems are commonly used on ships and airplanes for detection of obstacles interfering with their intended pathways. The platforms used for radar imaging of natural resources are mostly satellites or airplanes as these offer large coverages at high resolutions, also at large distances.

### 7.1.1. The radar equation

The principle of radar is that an electromagnetic signal is transmitted by an antenna into the surrounding medium, where it is reflected by objects and some of the scattered energy is reflected back to a receiving antenna. In case of the same transmitting and receiving antenna, the radar system is described as “monostatic”, and when one antenna is transmitting and another receiving, the radar system is described as “bistatic”.

Understanding the radar concept and its different components facilitates the interpretation of the acquired radar data, commonly known as images. The different components can be considered in the so called radar equation, which is similar to the ALS equation, originating from the same theoretical concept. The radar equation is based on the ratio of the transmitted power \( (P_t) \) and the received power \( (P_r) \), stemming from the waves scattered by a target\(^1\), and can be written as:

\[
P_r = \frac{P_t G_t \sigma A_r 1}{4\pi R_t^2 4\pi R_r^2 L}
\]

where \( G_t \) is the gain of the transmitting antenna, \( R_t \) and \( R_r \) are the distances from the target to the transmitting and receiving antennas respectively, \( A_r \) is the effective area of the receiving antenna, \( L \) is system losses, and \( \sigma \) is the target’s radar cross-section (RCS). The RCS is thus the only parameter in the radar equation that is related to the target, and depends on the target’s physical properties such as size, shape and material. For a perfectly conducting spherical target, the RCS would be equal to the cross-sectional area, \( \pi l^2 \), assuming \( l \) to be the radius of the sphere and with the premise \( l \gg \lambda \).

While this holds for targets like single trees, remote sensing is often more concerned with entire forests, where the radar system integrates series of radar signals to create 2D images. In this case a more useful measure of the RCS is the radar scattering coefficient sigma nought, \( \sigma^0 \) [m\(^2\)/m\(^2\)], which is defined as the average RCS per unit ground area of the target, and in case of the same transmitting and receiving antenna, it is denoted as the

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\(^1\) Floyd M. Henderson (Editor) 1998, page 132
backscattering coefficient. It depends on the physical and electrical properties of the illuminated object, and on the wavelength, polarization, and incidence angle used by the radar system. It also depends on the local surface slope towards the sensor.

7.1.2. Microwaves

Remote sensing can be performed with electromagnetic waves at a wide spectrum of frequencies. Electromagnetic radiation at frequencies 300 MHz to 300 GHz are generally considered microwaves, where “micro” represents “small waves compared to radio waves”. The radar bands include frequencies from the entire range 3 MHz – 300 GHz, including also the HF and VHF bands in the radio frequency domain, which sometimes also are used for imaging radar. However, there is not one single definition of the radar bands, but one commonly used list which is based on the bands defined during World War 2, and this one has mainly been adopted to an IEEE definition. Additionally, each country usually add complementing definitions suitable for their purposes. Common radar bands are listed in Table 7.1.

Table 7.1. A list of common frequency bands and a brief description of the origin of their names. Adapted from Anon (2003) and Radio spectrum (2014).

<table>
<thead>
<tr>
<th>Band name</th>
<th>Frequency range</th>
<th>Wavelength range</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>VHF</td>
<td>30-300 MHz</td>
<td>1-10 m</td>
<td>Very High Frequency</td>
</tr>
<tr>
<td>P</td>
<td>216-450 MHz</td>
<td>0.7-1.4 m</td>
<td>P for “previous”</td>
</tr>
<tr>
<td>L</td>
<td>1-2 GHz</td>
<td>15-30 cm</td>
<td>L for “long” wave</td>
</tr>
<tr>
<td>S</td>
<td>2-4 GHz</td>
<td>7.5-15 cm</td>
<td>S for “short” wave</td>
</tr>
<tr>
<td>C</td>
<td>4-8 GHz</td>
<td>3.75-7.5 cm</td>
<td>C for “compromise”</td>
</tr>
<tr>
<td>X</td>
<td>8-12 GHz</td>
<td>2.5-3.75 cm</td>
<td>X for crosshair</td>
</tr>
<tr>
<td>Ku</td>
<td>12-18 GHz</td>
<td>1.7-2.5 cm</td>
<td>Ku for “kurz-unten”</td>
</tr>
<tr>
<td>K</td>
<td>18-27 GHz</td>
<td>1.1-1.7 cm</td>
<td>German “kurz” (short)</td>
</tr>
<tr>
<td>Ka</td>
<td>27-40 GHz</td>
<td>0.75-1.1 cm</td>
<td>Ka for “kurz-above”</td>
</tr>
</tbody>
</table>

Figure 7.1. The electromagnetic spectrum. Atmospheric opacity is shown along the top. ORIGINAL SOURCE: Fawwaz T. Ulaby and David G. Long, Microwave Radar and Radiometric Remote Sensing, University of Michigan Press, Ann Arbor, Mich., 2014. With permission of the authors. This review source: https://www.nap.edu/read/21729/chapter/3
One reason for using microwaves is their suitability to penetrate the atmosphere, especially at lower frequencies (Figure 7.1). For frequencies up to about 20 GHz microwaves generally penetrate also clouds and rain (Figure 7.2). Rain has a greater influence on microwaves than clouds, starting its attenuation effect at about 3 GHz ($\lambda = 10$ cm) and increasing with frequency. However, rain only start to have an important effect at frequencies exceeding ~15 GHz ($\lambda = 2$ cm).

![Figure 7.2. Rain (a) and cloud (b) attenuation at microwave and millimeter-wave frequencies. Source: (Richards, Scheer, & Holm, 2010)](image-url)

The penetration behavior of microwaves is also considered as an advantage when imagery of forests are concerned. Optical waves are mainly reflected at the top of the canopy, while microwaves can often penetrate the canopy, with the attenuation (weakening of the signal) being dependent on wavelength, moisture content and the vegetation density.
Figure 7.3. Illustration of the scattering (penetration) in a forest canopy with common remote sensing radar bands. Optical sensors using sunlight are included for comparison.

### 7.1.3. Electromagnetic waves in the radar context

The field of remote sensing is normally considering radiation and especially electromagnetic radiation. While Newton considered light as particles, James Clerk Maxwell showed that light can be described as electromagnetic waves with Maxwell’s equations. The processing and interpretation of acquired radar images is facilitated by a further understanding of electromagnetic waves.

An electromagnetic wave represents the temporal and spatial variations of an electric and a magnetic field in space, which is characterized by amplitude and phase. This can mathematically be represented by a complex number on the form

\[ S = Ae^{j\phi} \]

where \( A \) is the amplitude of the wave (expresses the power) and \( \phi \) is the phase (a value on the 0-2\( \pi \) interval, defined as \( \arctan(\text{Im}(S)/\text{Re}(S)) \)). The phase is related to the wavelength and is proportional to the distance traveled by the wave, which in the radar case corresponds to the two-way path between the transmitter and the object(s) and can be written as

\[ \phi = \frac{2\pi}{\lambda} 2R = \frac{4\pi R}{\lambda} \]

Figure 7.4. Illustration of a phase-zero transmitted sinusoid, which after travelling the distance 2\( R \) from the satellite shows a certain phase \( 4\pi R/\lambda \). Adapted from Ferretti et al. (2007), Fig. 1-5.
Each signal can be represented both in the time and space domain and the frequency domain and the relation is given by the Fourier transform, defined as

\[ S(f) = \int_{-\infty}^{\infty} s(t)e^{-j2\pi ft} dt \]

where \( S(f) \) represents the signal in the frequency domain and \( s(t) \) the signal in the time domain. The Fourier transform of a rectangular pulse of length \( \tau \) is a sinc function where \( \text{sinc}(x) = \sin(x)/x \). The difference between the highest and the lowest frequency components is called the bandwidth.

Figure 7.5. Illustration of how pulse length \( \tau \) and bandwidth \( B \) are related.

The frequency representation of the signal \( S(f) \) entails that each signal can be written as the sum of its individual frequency components, and by increasing the number of frequencies transmitted, a larger bandwidth is achieved. In other words, by frequency modulating the pulse (also called chirp signal), a larger bandwidth \( B \) of the transmitted pulse is achieved and an increased range resolution is the outcome (Figure 7.5).

Figure 7.6. Illustration of a baseband chirp signal in the time and frequency domains. Maximum of the Fourier transform magnitude has been normalized to unity.
7.1.4. Combination of waves

What happens if two or more electromagnetic waves are combined? This concept is called superposition of waves.

Interference

When two or more waves are combined (the waves are passing the same position in space at the same time), the total amplitude is given by the sum of the individual amplitudes. Constructive interference happens when the sum of amplitudes is larger than the individual contributions. Correspondingly, destructive interference happens when the sum of amplitudes is smaller than the individual contributions.

![Figure 7.7. Constructive and destructive interference. The black wave illustrates the combination of the red and blue waves.](image)

Coherence

Interference requires identical frequency of the involved waves, which means that their phase differences remain constant over time. Waves complying with this criterion are called coherent waves. The vector illustrations above can depict interfering waves when they are coherent (their rotational speed is constant). However, when absolute coherence cannot be achieved, coherence is considered as a measure of predictability. With higher coherence, the properties of the complementary wave are easier to predict, given the other one. The coherence can be described by evaluating the cross-correlation, which in the practical case of interferometry simplifies to:

\[
\hat{\gamma} = \frac{E[s_1s_2^*]}{\sqrt{E[|s_1|^2]E[|s_2|^2]}}
\]

where \(\hat{\gamma}\) is the coherence, \(s_1\) and \(s_2\) are the master and slave images, and \(E[\cdot]\) are the expectation values. However, the expectation value is an unknown and is estimated via spatial averaging. The previous equation simplifies to

\[
\hat{\gamma} = \frac{\langle s_1s_2^* \rangle}{\sqrt{\langle |s_1|^2 \rangle \langle |s_2|^2 \rangle}}
\]

\(^2\) (Bamler & Hartl, 1998; Ferretti et al., 2007; Lee & Pottier, 2009)
which can be interpreted as taking the average of an area with pixels containing the phase difference of the images \( s_1 \) and \( s_2 \), and to normalize it, this average is divided by the magnitude of the total coherence of the same pixels.

### 7.1.5. Polarization

An electromagnetic wave’s polarization is describing the motion and orientation of the electric field vector \( \mathbf{E} \). For example, a wave travelling in the \(+z\) direction in a Cartesian (x-y-z) coordinate system must have the direction of \( \mathbf{E} \) in the x-y plane. This means, that the electric field is oriented along both the x- and y-directions, and its amplitudes will each vary sinusoidally. Perpendicular to \( \mathbf{E} \), the magnetic field \( \mathbf{H} \) varies sinusoidally as well. The wavelength is measured in the direction that the wave propagates. The transmitted polarization is determined by the physical structure of the antenna and by its orientation. Linear polarizations are most commonly used in satellite radar systems, but both circular and elliptical polarizations can occur as well.

![Illustration of an electromagnetic wave](image)

Figure 7.8. Illustration of an electromagnetic wave (Chaisson & McMillan, 2003) The \( \mathbf{E} \) field is illustrated in the x-direction, the \( \mathbf{H} \) field in the y-direction, and the wave propagates in the z-direction.
7.1.6. Scattering

The definition of the backscattering coefficient $\sigma^0$ is the RCS per unit area, which can be considered as the sum of the individual scatterers within a resolution cell. This means that when lots of targets within a resolution cell reflect the EM wave at different distances (=phases) from the radar, all reflections will add up coherently to the complex sum of contributions from the individual targets (superposition principle, Figures 7.7 and 7.10). Some contributions will cancel each other out while some will add up, and the coherent sum becomes random on the $2\pi$ (or $–\pi..\pi$) interval. The result of this adding and extinction of phases is called speckle and will appear as “salt and pepper” in the radar images. The random phases originate from the distances between the antenna and the individual scatterers being uniformly randomly distributed on the scale of the radar wavelength (c.f. Smith 1998). The occurrence of speckle causes the amplitude measured from a homogenous area to be Rayleigh distributed, irrespective of the target.

Figure 7.10. a: Illustration of how individual scatterers add up to a coherent sum. b: Illustration of how many different scatterers contribute to the total scattering reflection within a resolution cell.
7.1.7. Geometrical effects

The quality of radar images can be affected by the acquisition geometry. The radar “sees” only the line-of-sight, which is called slant range. The radar image is acquired in this geometry, but naturally a “ground range” image is more useful, as it has the Earth as reference and depicts the objects similar to other geographic data products. Images in slant range will appear compressed in the range direction (Figure 7.11). By knowing the height of the acquisition platform and the incidence angle, correct ground range ortho corrected images can be computed by simple geometry; the ground range is simply the projection of the slant range, which corresponds to the slant range divided by the cosine of the incidence angle.

The acquisition geometry might even lead to some information losses, especially in hilly terrain. This is illustrated in Figure 7.14, which shows how the wave front reaches different parts of the ground at different times. In the illustration, the left most parts would normally be expected to be depicted first, but as the actual distance to the top of hill B is shorter than to the base of hill B, this radar echo will return before the echoes from the foot of the hill. This is called layover. Foreshortening is another geometric effect that can occur. This happens when the radar beam reaches the base of a tall feature tilted towards the radar (e.g., mountain A) before it reaches the top. This has the consequence that slopes become compressed in the range direction. Foreshortening and layover are two consequences which result from relief displacement. Figure 7.12 shows how the hills B and C are all affected by severe shadowing as well, which means that no radar beams reach the ground and information loss occurs, making it impossible to correctly reconstruct the ground even with advanced signal processing.

![Figure 7.11. Left: Illustration of how a slant range image translates to a ground range image. Right: Slant range images will appear compressed in the range direction.](image)
7.1.8. Resolution

The backscattered energy can be measured and the distance $R$ to the scattering objects can be determined as

$$ R = \frac{cT}{2} $$

where $T$ is the time between the transmission and reception of the radio pulse. The resolution in range $\Delta R$ is a measure of the minimum range difference of two objects still resolvable as two single objects and is determined by the length of the (matched filtered) electromagnetic (EM) pulse. The pulse length is inversely related to its bandwidth $B$ and the range resolution, $\Delta R_r$, can then be written as

$$ \Delta R_r \approx \frac{c}{2B} \approx \delta_r $$

An important property of this expression is that the range resolution is independent of range, which makes it ideal for spaceborne platforms as objects at very long distances can be resolved. This also explains why side-looking configurations are preferred over nadir-looking; the former implies that the returns from different scattering objects can be separated using the differences in time, compared to nadir images from which all objects reflect the transmitted EM pulse almost simultaneously (Figure 7.13).
Figure 7.13. Illustration of a) nadir-looking and b) side-looking configuration. In a) the returns from A and B are ambiguous while the ones in b are not, because of the separation in time.

The antenna pattern expresses the radiation pattern of an antenna where the main lobe of an antenna (=antenna beam width $\theta$) is defined as the 3 dB angle; that is to say that the angular interval having as extremes half of the power with respect to the look direction (Figure 7.14). The antenna beam width is directly proportional to the wavelength and inversely proportional to the antenna length $L$. This can be described as

$$\theta = \frac{\lambda}{L}$$

when $L \gg \lambda$, and is used in the derivation of the azimuth resolution (also called along-track or cross-range resolution).

The azimuth resolution ($\delta_{az}$) is given by

$$\delta_{az} \approx R\theta_{az} \approx \frac{R\lambda}{L_{az}}$$

where $\theta_{az}$ is the beam width and $L_{az}$ is the antenna length (also denoted aperture length) in the azimuth direction. This holds for real-aperture radars, and the azimuth resolution can then only be improved by increasing the length of the antenna or decreasing the wavelength. Because of the dependence on range, this is not a good solution for remote sensing from long distances and as will be shown in section 0, another solution is used for the spaceborne case.
Figure 7.14. Illustration of the 3 dB angle corresponding to the antenna beam width $\theta$ (≈22.5° in this illustration) of imaging radar.

### 7.2. Synthetic aperture radar (SAR)

To retrieve a 2D radar image with high resolution in both azimuth and range direction, a system with high bandwidth and long antenna is needed. To get a reasonable azimuth resolution of a couple of meters or even tens of meters from a spaceborne platform, the antenna needs to be several km long! This is in practice impossible and during the early 1950s the idea of synthesizing a large antenna by moving a smaller antenna along a path emerged and Synthetic Aperture Radar (SAR) was invented.

When either the target or the antenna is in motion along the range direction during the acquisition, this can be measured through the rate of change of the phase, which is commonly denoted as the Doppler frequency. The non-relativistic Doppler-frequency shift $f_D$ is given by Ulaby et al. (1982):

$$f_D = -\frac{2dR}{\lambda dt} = -\frac{2v_r}{\lambda}$$

where $v_r$ is the velocity between the target and the radar in the range direction and the other variables have the same definition as previously described.

Consider an antenna mounted on a flying platform, illuminating the ground with the azimuth beam width $\theta_{az}$ (Figure 7.15). An object will then start being illuminated from the transmitted beam as the front of the beam reaches the object. While the flying platform is moving forward, the object will remain within the beam as the platform moves until the rear of the beam is passing the object. The beam is constantly created, as new short radar pulses (chirps) are transmitted over and over at high frequencies. The

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frequency of the transmitted pulses is called pulse repetition frequency (PRF), and is often on the order of kHz. Each transmitted pulse will bounce against the object and be reflected back to the radar, but as the radar is constantly moving, each pulse will be affected by a different Doppler frequency shift (Doppler Effect, which is the same as can be experienced from the siren of a passing ambulance). The pulse to pulse Doppler shifts can be calculated and by using signal processing, all transmitted pulses that hit the object can be combined coherently. This means that the antenna constituted by the synthetic aperture corresponds to the ground projected length of the azimuth beam width, which thereby allows for a high azimuth resolution as well.

![Image](image.png)

Figure 7.15. Illustration of the SAR principle for the modes ScanSAR, stripmap (SM) and spotlight (SL) for TerraSAR-X.

### 7.2.1. Orbits

To denote the direction of a satellite, it is put in relation to the globe with the North Pole at the top and the South pole at the bottom. When the satellite is depicting the region of interest from below against the north, it is called ascending. When the satellite is going against the South pole, it is descending.

![Image](image.png)

Figure 7.16. Illustration of satellite orbit directions.
7.2.2. Spatial resolution
The azimuth resolution can be described as the capacity of distinguishing between different Doppler frequencies of two neighboring targets along-track. This means that the Doppler bandwidth \( B_D \) is given by

\[
B_D = \left| f_{D,\text{max}} - f_{D,\text{min}} \right| = 2\left| f_{D,\text{max}} \right|
\]

which is found at the endpoints of the synthetic aperture, if the antenna is aligned against zero-Doppler direction. The diffraction limited resolution is given by Eq. (5), where \( L_{az} \) is the antenna length in the azimuth direction. The approximation is valid for \( L \gg \lambda \) and hence the Doppler bandwidth can be approximated as

\[
B_D = 2\left| f_{D,\text{max}} \right| \approx \frac{2\theta v}{\lambda} \approx \frac{2v}{L_{az}}
\]

where \( v \) is the antenna’s orbital speed, which in turn gives the azimuth resolution as the time-resolution multiplied by the platform radial velocity. This is approximately the same as the reciprocal of the Doppler bandwidth, which gives

\[
\delta_{az} \approx \frac{v}{B_D} \approx \frac{L_{az}}{2}
\]

In contrast to real-aperture radar (RAR) systems, the azimuth resolution of a SAR system gets better with a smaller antenna. This is, however, only valid when \( L \gg \lambda \) and for a system where the integration angle is limited by the antenna beam width.

The SAR range resolution \( \delta_r \) (derived in Eq. X) remains the same as in the RAR system, with the bandwidth \( B_D \) as the major limitation of \( \delta_r \). In practice, to avoid ambiguities, there is a need of using a higher pulse repetition frequency than the Doppler bandwidth \( B_D \). This translates to an upper limit of the range swath size and also a lower limit on the azimuth antenna size. The lower limit is proportional to the wavelength, so that for long wavelengths large antennas are needed. These constraints by range and azimuth ambiguities hence imply that the swath width and the azimuth resolution cannot be adjusted independently; high azimuth resolution gives a narrow swath and vice versa\(^4\). The derivation in this section is valid for stripmap (SM) mode and not ScanSAR or spotlight (SL) mode.

7.3. Radar signal processing

7.3.1. Radargrammetry
The term radargrammetry can be deduced from the optical term stereogrammetry but applied to radar images. Its fundamental principle is the stereoscopic viewing of an object from somewhat different angles, which can be related to the perception of depth vision by humans. By

\(^4\) Ulaby et al., 1982
combining disparities and convergences, stereo imagery can be attained in our brains. Due to the geometric and radiometric properties of SAR images, which differ from those of optical images, more time is generally required for the eyes to adjust to stereo viewing of SAR images in order to perform visual interpretation. However, depth perception is an active process and object recognition can be trained over time.

In radargrammetry the disparity principle is used to compute the terrain elevation from the measured parallaxes between two images, acquired at different angles\(^5\). The images are in contrast to InSAR or Polarimetric InSAR (but in similarity to optical photogrammetry) entirely relying on the backscatter intensity of the SAR images. The intensity \(I\), is defined as the amplitude squared, \(A^2\), and produces an image similar to a black and white photograph.

During the 1980s, the SAR systems improved and both same-side and opposite-side stereo viewing were later demonstrated, where the latter was found superior to same-side stereo\(^6\). However, the opposite-side configuration brought severe illumination difficulties, as they got so distinct that despite the geometrical advantages, stereo-viewing became so difficult that the success rate in finding corresponding points and features was limited. An illustration of different system configurations of radargrammetry is shown in Figure 7.17.

Since new launches of different satellite sensors tend to lead to hype over radargrammetry, and in turn lead to research with uneven steps over the years, it should be stressed that same-side configurations have been preferred and is the primary research path since the late nineties. In any case, the sensitivity of stereo measurements is increased with increased intersection angles, as the stereo exaggeration factor increases with larger observed parallaxes. This means that larger intersection angles technically give an increased height accuracy of the extracted terrain elevation. On the other hand, as has been discussed, the geometric similarities are important for an appropriate object recognition, which obviously is facilitated by small intersection angles as this gives more identical images. The compromise has often been to use intersection angles in the range of 7 to 25° and with small temporal differences for the acquisitions, in order to attain suitable geometric parallaxes and smaller radiometric disparities\(^7\).

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5. Thierry Toutin & Gray, 2000
6. Fullerton et al. 1986; Toutin 1996
7.3.2. Interferometry

SAR interferometry is a remote sensing technique that enables accurate measurements of geophysical parameters, which mainly are related to height changes. That could, for example, be subsidence, glacier movements and forest heights. The basic concept of radar interferometry, is to compare for a given scene, the phase of two or more complex radar images, acquired from different locations or at different times. Acquisitions from different locations can be achieved by acquiring data from different orbits, or like the Shuttle Radar Topography Mission (SRTM), by attaching a 60 m long mast to the space shuttle.

Currently (2016), the German space agency (DLR) operates the mission TanDEM-X, where two satellites are flying in a close formation. Each satellite can send and receive data, with the other satellite receiving the same signal, from the slightly different location. When a satellite setup with only one satellite is observing the same area from the same orbit but exploiting repetitive acquisitions from different times, the term temporal baseline can be used. SAR interferometry combines the images acquired at different locations or times, depending on the type of interferometry performed\(^8\). When a temporal baseline is used, changes that take place

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8 Bamler & Hartl, 1998
between the two acquisitions can be found. This is useful for measurements of, for example, land subsidence or glacier movements, which often take place at a rather slow pace. Forest also tend to change at a rather moderate speed. However, because of the non-solid character of a forest canopy, the complex geometric properties of trees cause incoherence, and attenuation of the radar signal, which make interferometry impossible. Hence, acquisitions from different locations at the same time, for example, from SRTM or TanDEM-X, are highly preferred when forest applications are considered.

**Technical aspects**

The phase of each SAR pixel contains information about the range, with an accuracy corresponding to a small fraction of the radar wavelength. This makes it possible to detect as small path length differences as centimeters or millimeters, which can be the case for subsidence. The accuracy is independent of the distance between sensor and the scene, which makes SAR interferometry suitable both for airborne and satellite borne remote sensing. However, a fundamental challenge of SAR interferometry is that the measured range difference is ambiguous with the wavelength. The resolving of this ambiguity is called phase unwrapping, and is usually dependent on some external data, for example, a rough height model of the scene to be unwrapped.

The one-way range difference $\Delta r$ is proportional to the height difference $\Delta h$, with the distance $r_0$ being large and the vertical baseline $B_\perp$ being small.

$$\Delta r = \frac{B_\perp}{r_0 \sin \theta_i} \cdot \Delta h$$

where $r_0$ is the slant range, $\theta_i$ is the local incident angle, and $B_\perp$ is the baseline perpendicular to the line of sight. In a coherent radar, this range difference $\Delta r$ corresponds to a measured phase difference $\Delta \phi$

$$\Delta \phi = m \frac{2\pi}{\lambda} \cdot \Delta r$$

where $\lambda$ is the carrier wavelength. The factor $m$ describes the range difference as either just the receive path, or both the transmit and the receive paths. Therefore, $m$ is equal to one for a single-pass SAR interferometer where one antenna transmits and (at least) two antennas receive, and $m$ equals to two for a repeat-pass system, where merely the same antenna is both transmitting and receiving.
Figure 7.18. Across-track SAR interferometry employs antennas moving parallel but mutually displaced from one another. The flight direction of the satellites is perpendicular to the image (out of the page). The slant range $r_0$, the incident angle $\theta_i$ and the perpendicular baseline $B_\perp$ are all defined in a plane perpendicular to the flight paths. A change in surface height by $\Delta h$ causes a change in range difference by $\Delta r$.\(^9\)

**Steps in interferometric processing**

Interferometric processing is here described as the process of generating a digital elevation model (DEM), from suitable SAR acquisitions. To perform SAR interferometry, the two involved images have to be aligned, such that the corresponding pixels in respective image reflect the same features on ground. This alignment is called co-registration and interferometry requires sub-pixel accuracy of this step. The first image is multiplied with the complex conjugate of the second, to form a so called interferogram. The complex conjugate means taking the phase with opposite sign, in order to obtain the phase difference between the images. The interferogram is a complex image, describing the complex coherence, usually denoted $\tilde{\gamma}$, possessing two-dimensional values, which makes it possible to derive the phase $\varphi$ as $\arg(\tilde{\gamma})$ and the (non-complex) coherence $\gamma$ as the magnitude of the interferometric coherence, $|\tilde{\gamma}|$. The coherence $\gamma$ describes the degree of correlation between the two radar images and is hence the main limitation of the accuracy of the phase measurements. There are different reasons for decorrelation, but in order to suppress this phase noise, spatial averaging (denoted multilooking) is usually applied before extracting the phase values from the interferogram.

By inserting the first equation into the second above, the following relation can be found:

$$\frac{\Delta \varphi}{\Delta h} = \frac{2m\pi B_\perp}{\lambda r_0 \sin \theta_i}$$

This equation describes the sensitivity of the interferometer to small height differences $\Delta h$. From this relation, it can be noted, that by increasing the

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\(^9\) Krieger, Hajnsek, Papathanassiou, Younis, & Moreira, 2010)
perpendicular baseline $B_\perp$, the sensitivity for height changes also increases. However, with increased baselines, the difference in appearance of the scattered resolution cells also increases, which means that the correlation and coherence decreases, until it finally vanishes. The spatial decorrelation can be removed by a process recognized as range filtering, at the cost of decreased range resolution. With a larger bandwidth of the SAR system (which most modern SAR systems have), this problem decreases.

Another, more important restriction of the useful maximum baseline, results from the ambiguities in the phase to height conversion. The interferometric system measures only relative heights, ambiguous by an integer $m$ times $2\pi$. This height of ambiguity (HOA), can hence be expressed as

$$\Delta h = \frac{\lambda_0 \sin \theta_i}{2B_\perp} \cdot m \cdot 2\pi$$

for the bi-static system currently considered. When a monostatic system is used, another factor $\frac{1}{2}$ has to be included.

The relation between the baseline, the perpendicular baseline (two x the effective baseline) and the HOA is illustrated in Figure 7.19.

Figure 7.19. Relation between baseline and HOA

The final step in the DEM generation is geocoding of the height image. This is transforming the height pixels from the radar geometry (slant range) to a map geometry, for example the UTM projected on some ellipsoid. This transformation can be done by creating a lookup-table (LUT), which maps each pixel in the radar geometry to corresponding pixels in the map geometry. This LUT can hereafter also be used to geocode for example intensity and coherence images to the map geometry.

### 7.3.3. SAR Polarimetry

The electromagnetic waves transmitted by a SAR system possess a certain polarization. As the waves interact with objects on ground, some of these can change polarization, dependent on the interactive object geometries. Most often, the same polarization is transmitted by a SAR system as received, e.g., the antenna transmits horizontal waves and received horizontal waves. Radargrammetry and interferometry work well with a single polarization, preferably the vertical-vertical (VV) or horizontal-horizontal (HH), as these polarizations tend to be stronger then the cross-polarization horizontal-vertical (HV) or reversed (VH). However, some
SAR systems can acquire fully polarimetric data, which contains information about both HH, VV and HV/VH (VH bears the same information as HV). The full polarimetric data is acquired to the cost of lower resolution, as parts of the digital antenna is sacrificed for the additional polarizations. The advantages, is that the scattered properties can be fully described, using a so called scattering matrix, e.g. the Sinclair matrix or Jones matrix. After extensive mathematical operations, the data can be efficiently used for classifications of for example vegetation, buildings or water.

Figure 7.20. A classified AIRSAR L-band image of the frequently used San Francisco bay area, illustrated in natural colors. Data collected by JPL-NASA-Caltech. 2 looks multilooking, resolution 10m x 10m.

7.3.4. Polarimetric SAR interferometry

By applying interferometric principles and theories to multi-polarized data, polarimetric SAR interferometry (Pol-InSAR) can be achieved. Many current forest related theoretical models originates in Pol-InSAR, such as the interferometric water cloud model (IWCM), random volume over ground (RVoG), or the two-level model (TLM).

Self Study Questions
Further Reading
9. METHODS FOR ANALYSIS OF REMOTE SENSING DATA

**Classification.** Analysis methods for assigning the values of remote sensing data into specific categories or classes.

**Supervised Classification.** An analysis approach used when training data are available, and used to develop the model used for classification or prediction.

**Estimation.** Analysis methods used for obtaining continuous variables.

**Change detection.** Method for detection changes by comparing remote sensing data from more than one timepoint.

Creating new information from remotely sensed data is performed by using the values of the remote sensing data in a statistical model which then interprets the data into information relevant for the user. The choice of analysis method often depends on the desired outcome (thematic or continuous values) and the characteristics of the input data. The different models or algorithms involved in the analysis method have their own characteristics, with associated advantages and drawbacks. The method of analysis chosen is often a subject for research in remote sensing.

### 9.1. Classification and estimation

We can categorize the approaches used to produce new information from remote sensing data in a number of ways, but perhaps the most general way is to categorize them by whether the end product will be thematic (classes) or continuous values. Classification algorithms are used to produce thematic classes, while estimation methods are used to produce continuous values.

In general, *classification* refers to the assigning of any group of things into their own categories, based on the data characteristics. In remote sensing, it refers to assigning the values of remote sensing data into specific categories or classes. It is an exercise in pattern recognition, and a number of analysis methods used to categorize remote sensing data come from the field of pattern recognition.

There are several types of classification methods, which use a statistical basis for class assignment of the pixels. Some of these will be discussed in the following sections of this chapter. There is no single correct classification method to use. However, some methods may be better than others depending on the goal of your project and the available data. Methods are continually being developed and tested.
Estimation methods are used for obtaining continuous variables, and this approach is commonly used for forest mapping (for example stem volume, age, or species proportions). Estimation of continuous forest variables is a particularly common method with laser data, radar data, and 3D aerial photo models. Examples of estimation methods are kNN, regression analysis, regression trees, or random forests. A more strict definition is that estimation is used for statistical methods that establish a relationship between variables, e.g. an estimation of a regression functions that gives the relation between a dependent variable (e.g. stem volume) and on or several independent variables (e.g. spectral bands in a satellite image and/or metrics from 3D point clouds). The process of applying that estimated function on all forest pixel is then called prediction in some texts, but estimation in other texts. Some of the methods used for establishing relationships between remote sensing data and forest variables, for example Neural Networks comes from other science fields than statistics, for example computer science, still, we often use the word estimation for all these techniques.

9.2. Background information

9.2.1. Use of reference data in classification or estimation

Before going into analysis methods, some words about reference data are needed. Reference data can be taken from field visits, aerial photo-interpretation, or ground-based inventories, for example. The term “ground truth” is sometimes used to refer to reference data. Reference data have assigned or measured variables describing the properties of the sample plot or an area (e.g., land cover class, percent shrub cover, or basal area) and often have associated geographic coordinates. The common geographic locations allow associating the remote sensing data with the variables from the reference data (see Figure 9.1). More information on reference data is given in Chapter 7.

Figure 9.1. Example of a 10 m radius NFI plot corresponding to the same geographic location in Landsat TM data (25 m pixels).

Spectral data classes and Information classes

When classifying spectral data, there are some terms to be familiar with. These are spectral classes and information classes. The classes in your classification scheme that you want to identify are called information classes. A single information class may be represented one or many spectral classes. This is demonstrated below in Figure 9.2.
Figure 9.2. On the left the information class “Coniferous forest” is shown although you can see two distinct spectral groups, which are shown on the right as the two spectral classes that belong to the “Coniferous forest” information class. This might be due to training data collected from, for example, different slope directions.

When using spectral data, we can say that there are two basic categories of classification methods:

- spectral pattern recognition
- spatial pattern recognition

The main type of classification method used is based on spectral pattern recognition principles, meaning that the pixelwise colours of the multispectral images are used to separate different vegetation types. In this chapter, we will primarily discuss spectral pattern recognition methods.

There is also spatial pattern recognition where the spatial relationship between image pixels are used. Examples of spatial measures are:

- image texture (important with high resolution imagery, with 2 m pixel size, or better)
- object shape (e.g., using segmentation)
- context in the landscape (e.g., a set of rules how to interpret distance to water is used).

### 9.3. Classification methods

Classification has previously been applied mainly with spectral data from satellite sensors. However, it’s important to realize that any data can be divided (with more or less success) into thematic classes.

We can further categorize classification methods into different groups, which in addition help us choose which method we want to apply. The first main categorization is based on whether we have sufficient reference data to use in the analysis. This division is
• Supervised classification
• Unsupervised classification

Yet another categorization of classification methods is based on the statistical properties of the algorithm. This division is

• Parametric methods
• Non-parametric methods

Supervised classification is where reference data are available and used directly to identify the spectral values in the satellite image belonging to that vegetation type. The reference data are used to identify a subset of the pixels in the image to build a model for the classification algorithm. The model is then applied to the entire satellite image. These reference data are referred to as “training data”, and a good training data set should assign class labels to the range of spectral values present within the satellite image. The result is a classified thematic map.

Unsupervised classification methods first cluster the remote sensing data automatically into a decided number of classes. The classification is made based on the statistics of the image, without use of reference data. Reference data are then used afterwards to assign a class label to each cluster. Unsupervised methods are often used when reference data are sparse or inadequate.

9.3.1. Unsupervised classification

Unsupervised classification can be divided into the following steps:

• Clustering (analysis)
• Assigning the clusters to information classes

Many different algorithms for this exist, for example, “K-means” clustering or the ISODATA algorithm. The following describes a typical process for unsupervised classification:

1) the user selects a number of classes or “clusters” based on the goals of the project and also the spectral properties of the image. For example, maybe only four clusters are needed if the information classes “water” and “other” are wanted (because water is so spectrally unique and easy to identify from most other land cover). However, if you want to have several information classes, you will most likely need to give a higher number of clusters than your number of information classes.
2) the computer first divides the multi-spectral satellite data into four clusters based on the mean values
3) each pixel in the image will belong to one of the clusters to which it has the spectrally closest class mean
4) new means are calculated for the classes according to the pixel values assigned to them
5) Procedures 3 and 4 are repeated until stable means are found.
6) Once the clustering algorithm has finished, the spectral classes in the output must then be manually assigned to the relevant information classes.

The following situations may occur when assigning an information class to a cluster:

- One spectral class will clearly belong to one information class (this is the easy case)
- Several spectral classes will belong to one information class (the spectral classes should then be labeled with the single information class)
- One spectral class will belong to several information classes (this means there will be problems obtaining an accurate map product based on spectral data only).

9.3.2. Supervised classification

Supervised classification is divided into the following steps, in this order:

- Training data set creation
- Classification (or analysis).

The training data creation stage is where a data set which associates real world vegetation characteristics with the remote sensing data, and the data set is then used to build a model. Individual training areas are identified and delineated (these can be single pixels or they can be polygons which enclose several to many pixels) in the image. Each one of these training areas is individually referred to as a training data sample, and a group of training samples is called a training data set. In a training data set, it is important that all spectral classes are included. In general, for each information class you will have several training data samples. The training data set can consist of all the training data samples for all the information classes in the image.

Training data samples can be obtained either automatically or manually. Automatically derived training data may be made when existing inventory data are used, where the remote sensing data are extracted from that coordinate. A manual way to obtain training data is by drawing a single point or polygon around a single pixel or group of spectrally similar pixels (Fig 9.2).

The aim of a training data set is to accurately represent the range of values in the remote sensing data that are associated with each vegetation type. For example, several training data samples for a particular vegetation class will have basic statistical properties such as a mean value and standard deviation. This describes the range of values that are typical for this class. The aim is to have distinct values for the different classes, and for that reason, training data samples should be spectrally similar and not have a very large standard deviation.

Training data can be selected subjectively or objectively. Subjectively selected training data is done by purposely identifying optimal sites for
developing a training data set. In other words, it is not based on a probability-sample and has been chosen subjectively by the analyst. This approach can lead to a well-behaved training data set, and (often) a better map product, and is allowed for training data. A disadvantage of this approach is that the analyst may be biased about which areas they choose.

In forestry, there is more of a tradition to carry out objective ground sampling. Therefore we mention that another source of training data can be, for example, NFI plots with GPS coordinates or other pre-existing inventory data. An advantage of this approach is that if the reference data plots are located in a systematic or random manner, proportions of classes can be estimated from proportions of plots. A disadvantage is that the sample may consist of many mixed (spectrally non-homogenous) pixels (Fig 9.3).

One cause of many misclassifications or difficulties in classification is the mixed pixel (sometimes called a “mixel”). Mixed pixels occur when there is more than one land cover type within a single pixel. This can occur especially when the pixel size is large, when the landscape is made up of small vegetation units, but in general, it can occur with any image just because it is impossible for a pixel to always fall evenly on clear boundaries of vegetation, nature doesn’t work that way. The problems occur because mixed pixels can have spectral signatures that imitate the spectral signature of another class, or else have their own unique signature. For example, look along the borders between stands in an image. This is a typical area for mixed pixels, and a typical problem area. Consider what happens when a pixel covers an area of half water and half forest.

Although a subjective approach is acceptable for training data creation, it is worth repeating that accuracy assessment should be always be with an objectively collected dataset to evaluate the result.

The classification or analysis stage where a class membership is assigned to each pixel. There are many different supervised classification algorithms which can be used, for example, Minimum distance to means Maximum Likelihood Classification, Decision Trees, Random Forest or Neural networks. These algorithms will be discussed more in detail later in this chapter. Often the classification process is an iterative one, where the training data and the remote sensing input data are adjusted, as well as the parameters of the classification method, until the result is satisfactory.

Post-classification smoothing is a post-processing method which aims to create a “smoother” looking map. The result from supervised classification may require smoothing because it often looks more “pixelly” or has a “salt-and-pepper” look than does the outcome from unsupervised classification. This technique is described further in the section on manipulation of images.
9.3.1.2. Parametric and non-parametric classification methods

There are many different supervised classification methods, which can be separated into two groups: parametric methods and non-parametric methods. Parametric methods can be used if the training data meets the assumption that they have a multi-variate (i.e., all bands) normal distribution (Fig 9.4). With non-parametric methods the above assumption is not necessary to meet.

Fig 9.4.

Parametric methods are useful when the characteristics of the data are less complex. Typical methods include linear discriminant analysis (LDA), and maximum likelihood classification. Non-parametric methods include decision and regression trees, random forest, kNN, support vector machines, and neural networks. The previously named non-parametric classifiers all fall under a category called “machine-learning” algorithms, in which the data are iteratively processed until the accuracy of the outcome is optimized.
While non-parametric methods do not require having training data that follow a normal distribution, the results can be affected by having a training data set which is too small, or has too many of a single class (an “imbalanced data set”). In general, non-parametric methods require a larger number of training data sets.

### 9.4. More on training data for supervised classification

There are a number of characteristics of the training data used for supervised classification that need to be considered. These include

- number of training data samples needed,
- size of the training sample in relationship to the phenomena in the landscape as well as the remote sensing data spatial properties,
- statistical properties of the sampling scheme,
- interaction between the classification method and characteristics of the training data set, and
- quality of the training data.

#### 9.4.1. Number of training samples

A sufficient number of training samples and their representativeness are critical for image classifications. However, precise determination of the number of training data samples needed to achieve an accurate classification is elusive. This is, in part due to the “catch-22” nature of needing detailed and accurate class and spectral information on which to base estimates of the number of training data samples needed. It is also due to the complex interaction of different factors that influence the classification results.

The number of samples necessary to obtain adequate representation of the spectral variability present in a class can be determined statistically by sampling. Traditional distance measures of separation between classes are also often used (e.g., Jeffries-Matusita distance or Transformed Divergence), but these may be poor predictors of actual classification accuracy as they don’t provide information about the adequacy of spectral representation of an individual class.

Not only the representation of a class’ spectral variability needs to be taken into consideration, but also the spectral similarity or dissimilarity to other classes. As an example, water has a spectral signature so distinct from many vegetation types, that a full description of the spectral variability of water may not be necessary for accurate classification. Two very spectrally similar classes, such as willow and mesic heath require careful and full assignment of their spectral characteristics if one hopes to accurately classify these overlapping classes.
Considering class spectral overlap, Van Niel et al., (2005) found that for each class, only 2 to 4 times the number of input variables were sometimes necessary. Foody used neural networks (Foody, 1999) and Support Vector Machines (Foody and Mathur 2006) which depend on good separation between class boundaries in variable space, therefore only needing to include minimum and maximum variable values for all classes, allowing the training data set size to be smaller.

9.4.2. **Interaction between the analysis algorithm and the training data**

The characteristics of the supervised classification method exert an influence on the requirements from the training data set. As an example, statistical classifiers using the mean vector for class assignment will be influenced less by outliers in the training data, while classifiers such as Neural Networks can be highly influenced by individual poor quality training data samples. Some classifiers, such as maximum likelihood, require a minimum of \( p+1 \) (\( p = \) number of input variables) training samples per class to build statistics (e.g., covariance matrices). Based on this, Mather (2004) recommended that 10 to 30 times \( p \) training samples per class should be used. Non-parametric methods don’t face this restriction, but are still affected by the total number and frequency of classes in the training data.

In the estimation of continuous values, a larger training data set is generally required. Mathys et al. (2009) found that for estimation of continuous parameters, a training data set consisting of the whole range (0-100%) of the parameter was necessary. In fact, training samples containing mixes or 0% of the parameter were more important than training samples with 100% representation of the parameter.

9.4.3. **Size of the training data sample**

The size of the training data sample plot used for training data is also important. Gong and Howarth (1990) suggested that training data were best selected using single pixels and a systematic sample, however Chen and Stow (2002) said this was more suitable for homogeneous land cover types. Chen and Stow (2002) tested training samples taken on single pixel level and in blocks of pixels, finding that training data set size mattered more when single pixel training samples were used, and that blocks of pixels produced higher accuracies for training in heterogeneous landscapes. However, their result may have been dependent on the urban land use classes particular to their study. When the landscape of a study area is complex and heterogeneous, selecting sufficient training samples becomes difficult (Lu and Weng 2007).

9.4.4. **Statistical properties of the sampling scheme**

The frequency of the classes as represented in the training data has an influence on the results. In large area projects, it is desirable to have *a priori* information on the frequency of classes in the area (Cihlar 2000), as input to the supervised classification. For this reason, prior probabilities are assigned in Bayesian classifiers. Additionally, weights can be added in Decision and Regression Tree models to counteract imbalanced data (Xie 2009). Rare
classes are often a desired class in map products, and sufficient training data may need to be collected specifically for this purpose.

9.4.5. Quality of the training data
Quality control and refining training data in order to obtain accurate classifications is important to do. Some researchers suggest reducing the effects of outliers in the training data by weighting training samples according to their quality or by subjecting training data to majority-vote classifiers to detect mislabeled data. In the ideal case, training data should be evaluated before used in the classification/estimation process. To evaluate training data, the following can be done:

- Looking at scatter plots of training data
- Using spectral distance measurements between training data sets (e.g., Transformed divergence, Jeffries-Matusita)
- Investigating the classification result and the effect of inclusion/exclusion of potentially problematic training data samples
- Usually a combination of all of these is done
- To improve the result, you may need to add and delete training data samples…

The training data set almost always needs be evaluated and changed to achieve optimal classification accuracy. This process can take a considerable amount of time!

There are also temporal aspects to recognize when using training data. Reference data may be collected from dates differing from the satellite imagery and this may cause erroneous class assignment. The timing of the image acquisition and the vegetation phenology must be considered in relation to the training data. In the case of the alpine landscape, the natural seasonal dynamics and change in moisture conditions that can occur within the growing season as well as from one year to another in the alpine region pose challenges to using training data and satellite data from different time points. Within managed forest landscapes, silvicultural activities such as thinning and clear-cutting need to be identified.

9.5. Algorithms for Supervised classification

9.5.1. Maximum Likelihood Classification (or Discriminant Analysis)
In remote sensing, practitioners often refer to the “Maximum Likelihood” (ML) supervised classification method (Lillesand et al., 2008). Maximum likelihood classification was often used for land cover mapping projects in the 1980’s and 90’s, and is still used today. In essence, when remote sensing practitioners refer to “maximum likelihood classification”, they are most
often using what statisticians refer to as Bayesian quadratic discriminant analysis (QDA). Bayesian QDA uses statistics about the prior probability of a class occurring, based on the frequency of that class in the training data. We go to the length of explaining this because, it would actually be more correct to call “Maximum Likelihood Classification” by the same term used by statisticians (Bayesian QDA), however, this has not been commonly done in the remote sensing literature. The following should be read with that in mind.

Discriminant analysis (and our “Maximum Likelihood Classification” method) is a method by which one assigns class membership to an unknown observation based on a sample of data with known class memberships. The example here is that our known data comes from our training data, and the unknown data are all the other pixels in the image.

From the training data, a probability density function based upon the training data, as well as prior probability weights for each class are used to calculate the probability of the unknown pixels values “belonging” to the values in the sample data set. Then a posterior probability that an observation belongs to a certain class is calculated using the prior probabilities, the probability density function, and the probability of occurrence for that observation.

The remote sensing term “Maximum Likelihood Classification” is derived from the fact that a maximum likelihood estimator rule is used in Bayesian discriminant analysis, in which the observations are assigned to the most “likely” (i.e., highest probability) class, in order to maximize correct classification assignment. An illustrative example of how the Maximum Likelihood Classification algorithm works is shown in Figure X.

Figure X will demonstrate the Maximum Likelihood algorithm.

Discriminant analysis works best following the assumption that the data for each class and variable are normally distributed (meaning that there is a Gaussian distribution of the spectral data values within the training data for a single land cover class (or whatever class you wish to map). This means that Discriminant analysis (i.e., our Maximum Likelihood classification) is a parametric method. In some cases, the covariance matrices of the different classes in the data may have the same distribution, and in this case, a linear discriminant analysis (LDA) can be used. In cases where covariance matrices are not the same between classes, quadratic discriminant analysis should be used.

In practice, the application of Maximum Likelihood classification in land cover mapping has often left out the use of prior probabilities of class occurrence due to lack of sufficient information about the frequency of class occurrence. When “maximum likelihood” classification without prior probabilities is used, then this is actually “Bayesian discriminant analysis with equal prior probabilities for all classes”. If equal probability is assumed and no weights are used in the training data, the result may be that more frequently occurring classes in the training data will be under-classified.
(errors of omission) in the resulting map and less frequently occurring classes will be over-classified in the map (errors of commission). Several remote sensing studies have pointed out the utility of including prior probabilities within the “maximum likelihood” classifier, finding that it improved land cover classification accuracy, particularly for spectrally similar classes.

The expression used for Maximum Likelihood classification is often based on Richards (1999), shown in Eq. X below.

\[ g_i(x) = \ln p(\omega_i) - \frac{1}{2} \ln |\Sigma_i| - \frac{1}{2} (x - m_i)^T \Sigma_i^{-1} (x - m_i) \]

where:

- \( i \) = class
- \( x \) = n-dimensional data (where \( n \) is the number of bands)
- \( p(\omega_i) \) = probability that class \( \omega_i \) occurs in the image and is assumed the same for all classes
- \( |\Sigma_i| \) = determinant of the covariance matrix of the data in class \( \omega_i \)
- \( \Sigma_i^{-1} \) = its inverse matrix
- \( m_i \) = mean vector

In practice, with Maximum Likelihood, two approaches with the training data can be used (as demonstrated in Fig 9.3).

- separate training samples can be merged into a single training data statistic for a single information class, or
- individual training samples can be used (i.e., spectral classes) and thereafter be recorded in the end product belonging to an information class (e.g., Coniferous 1, Coniferous 2, etc…).

Figure 9.3
An advantage (or potential disadvantage, depending on the data) of the latter approach is that the standard deviation of the training data will be smaller, and reduce the risk of including too many pixels in the satellite data (potentially not belonging to that class), resulting in misclassification. A disadvantage would be if the training data samples are too “narrow” having too low a standard deviation, to include the appropriate pixels. Your choice depends on the properties of your data.

Figure 9.4 is an ideal picture of a training set, and rarely are the classes so well separated. Figure 9.4 shows a training data set closer to reality.

![Reality of a training data set](image)

The training data set almost always needs be evaluated and changed to achieve optimal classification accuracy. This process can take a considerable amount of time!

9.5.2. CART and Random Forests Classification
Classification and Regression Tree (CART) methods have been increasing in use for land cover classification over the past decades. Three methods being widely used are decision trees, regression trees and random forests. Decision trees produce a categorical output, regression trees produce continuous variables, and the random forests algorithm is capable of producing both. These non-parametric methods have an important advantage over maximum likelihood classification in that data from different sources (e.g., spectral data, elevation derivatives, map data) can be combined, without the need for assumptions of normal distribution. These methods are also particularly useful when a large number of input variables are to be used.

9.5.2.1. Decision (or Classification) and regression trees
In decision trees, a hierarchical tree is constructed from the training data. The tree consists of root nodes, interior nodes and terminal or leaf nodes (Tso & Mather, 2009). Data splitting rules are constructed at each non-terminal node based on the training data’s spectral and class values. Splitting rules depend on the specific implementation of the decision tree, although most often they are based on determining the maximum information gain (Quinlan, 1993) and the lowest Gini impurity index (Breiman et al., 1984) based on the input variables at each node. In decision trees one variable is normally used for
splitting at each node, although multivariate decision trees have been developed (Friedl & Brodley, 1997). Pruning of the trees is often necessary to avoid over-fitting of the data, often accomplished by setting aside a portion of the training data to use for pruning. Regression trees also consist of the same node system, however univariate or multivariate regression functions are built to estimate continuous values.

9.5.2.2. Random Forests

Random Forests is a non-parametric method that can be used for both estimation and classification. It was developed by Leo Breiman, University of California, Berkeley and Adele Cutler at Utah State University. Note that forests do not have to do with the forest, but refers to the method used by a large number of classification (or regression) trees. Each tree gives an estimation or classification result, and the final result is calculated as a mean value (in the case estimate) or by "majority voting" (the classification). The advantages of the Random Forests is that one can use a large number of independent variables, including those that are correlated with each other, as well as different types of data (spectral data and map data) with different properties. Random Forests have effectively shown to give almost as good results as regression, without having to manually build a model of the relationship between independent and dependent data. Use of the random forests classifier has also produced classification results that are equally accurate or more accurate than other classification methods, and it is relatively robust to outliers and noise (Breiman, 2001).

Say that we are interested in finding forest the requires thinning. We believe that the stem/ha count is high and is within a certain height range, and we want to define such areas in a GIS to plan a control in the field. We create the classification tree in Figure 9.5 and use it in conjunction with vegetation ratio, \( V \), and the 90th height percentile, \( H_{90} \), in the form of breaks. Each question, or node, in the tree in a variable and can be answered with "yes" or "no". The tree flows into a number of end nodes, each of which corresponds to a class. In the example in the Figure, there are two classes: "no action" and "field check". A class can be found in multiple end nodes.

![Classification Tree Diagram](image-url)
Figure 9.5. Example of a simple classification tree that can be used to identify areas of dense forest \((V > a)\) within a certain height range \((b < H < c)\). Interesting areas frequented since the fields for the control of thinning needs.

In the regression case, the data set is not divided into categories as illustrated above, but instead constructed into trees so that each leaf contains a small, relatively homogeneous group of observations. The end node is a number (i.e., a continuous value) rather than a class. Training data, such as field inventory plots, are used to build the tree. The final result is calculated as the average number of the results from all the trees. The model obtained is then applied to all the “unknown” pixels in the image.

A tree is constructed by taking a so-called bootstrap sample of plots (it extracts a certain percentage of training samples which are used to internally quality check the model), by drawing with replacement. To build the first node it uses a user defined number of independent variables to test. The best variable that divides the sample clearly into two separate groups is selected. After that, each branch going out from the first node, leads to a new node which is constructed in the same manner as the first. The tree is considered complete when the number of sample plots in each leaf reaches a user-specified value.

There is a final assessment of the model quality which is given by the so-called OOB Error Estimate (OOB stands for “out-of-bag”). It is estimated by a given percentage of the sample plots that are not included in the training data that were used to create the Random Forest model. Random forests uses bagging (Breiman, 1996) as well as a random selection of the variables to consider at each node, therefore pruning of trees in random forests is not required.

Probability of class membership is based on the frequency of classes in the training data. Therefore, all the tree classification methods discussed here are subject to misclassifications due to imbalanced data (i.e., having an uneven distribution of training samples among the classes), and for this reason, weights and other modifications to the training data sets are sometimes made.

9.5.2.3. Support Vector Machines and Relevance Vector Machines

Both Support Vector Machines (SVM) and Relevance Vector Machines (RVM) are relatively new methods, which are based on so-called “kernel” methods. This means that they transform the original values into new values (go from the original feature space to a “kernel feature space” which has higher dimensionality and allows for a linear model to be applied). Support Vector Machines rely on a well-defined boundary (called the “hyperplane”) between two or more spectral classes, which is determined iteratively. Relevance Vector Machines are a Bayesian extension of SVMs (include prior probability measures). An advantage of SVM and RVM is that it does not require a large number of training data samples, but can work with many bands of input remote sensing data. A figure demonstrating SVM is shown in Figure 9.6.
9.5.2.4. Object-oriented image analysis (OBIA)

Note that classification can be carried out at the pixel level or at an aggregated level, such as within segments, which is referred to as an object-oriented image analysis.

9.6. Estimation algorithms

9.6.1. Regression

Regression is a statistical method for finding a relationship between two or more random variables. The method is model-based, i.e., it assumes that there is a special mathematical model that relates the two variables together. Although the method is based on a lot of theory, the basic idea is simple. Working with regression analysis is something of an art, but you can get far with fairly simple models. The following is a somewhat simplified description of regression.

The Figure 9.7 shows a scatter plot of field measured basal area average height, H, on the y-axis and the 90th height percentile, H90, on the x-axis.

![Figure 9.7. Scatter plots with 90th height percentile from laser data (X-axis) plotted against basal area weighted mean height (Y-axis). The arrows show how the regression function (solid black line) will result in use for elevation estimation.](image)

Each point represents a sample area where you have both field measurements and laser data. There tends to be a strong relationship between forest height and H90, which is evident in the scatter plot above, where you can see the points are well grouped along a line. By investigating the relationship between average height and H90, you can then estimate H in areas where field data is missing. Note that H90 is not the same as the average height of the forest, but only correlated with this! Correlation is a measure of the strength of a linear relationship between two variables, and the correlation coefficient can have values between -1 and 1. If y increases with x and the points lie exactly on a line, the correlation is 1, which means a perfect, positive linear relationship. If instead y decreases with x, and points lie exactly on a line, the connection is perfect negative correlation is -1. The
closer the correlation is to 0, the weaker the connection. In the data sets used in the Figure above, the correlation between H and H90 = 0.93, which means a very strong, positive relationship.

To begin with, you have to find the model that best fits the observed data. In the case of H and H90, it appears to be a straight line.

A Straight-line equation is written as

\[ y = k \cdot x + m, \quad (1) \]

where \( k \) is the slope of the line and \( m \) is the intercept where the line crosses the \( y \)-axis. The points on the graph are not exactly on a line, there is some random deviation or spread around the line. We write, therefore, the relationship between H and H90 as

\[ H_i = \alpha + \beta \cdot h_{90,i} + e_i, \quad (2) \]

where the index \( i \) denotes a certain field plot location and \( e \) is the deviation from the line at that pair of observations from field and remote sensing data. Using regression analysis, we can determine the values of \( \alpha \) and \( \beta \) that minimize the sum of squares of the distances \( e \), the so-called residuals, between the points and the line, under the constraint that the mean of all deviations is zero. Equation 2 describes H as a function of H90, and the variable H is called the dependent variable and the variable H90 is called the independent variable. H can also be called the response variable and H90 can be called the explanatory variable. The regression analysis also provides estimates of how strong the dependence is between the two variables. A common measure is the coefficient of determination. This quantity is often referred to as \( R^2 \) and indicates how much of the variance the regression model describes for the \( y \)-variable. The coefficient of determination is one of the tools usually used to select which variables should be included in the model. Another important measure of the model's accuracy is its spread around the regression line.

When a reliable function has been made, it can then be used to estimate the average height of trees in areas where there are laser data but no field measurements have been taken.

The above example addresses the simple linear regression. In many cases, several independent variables are used to describe the subject, and this is then called multiple linear regression. You may also need to transform the independent variables by squaring it, multiplying the two independent by each other, and so on. Some relationships are non-linear and can be linearized by logarithmic transformation prior to doing the regression. This is not addressed here, and those who want to go deeper into the theory are referred to the statistical literature.
Regression in practice
Since the estimation results generally are aggregated into stand averages, it is important with a good stand delineation. In Norway, stand deleniation and species assessment is done with manual interpretation in digital photogrammetric workstations, then other forest data are estimated on the basis of laser data. Stand delineation can also be made on the basis of segmentation of laser data (section 6.3). Manual editing may be needed and aerial photos are of course also useful in this work. Stand boundaries are stored in vector format in a GIS. Stands and sample plots are sometimes divided into strata for which separate regression functions are estimated. The stratification might for example be based on tree species, site index and age. These characteristics affect crown shape, which in turn affects the relationships between laser measurements and forest variables. Estimates can get higher accuracy if done for more homogeneous groups, but there are also examples of experiments where stratification has not improved outcome.

When the field data are collected, as well as possible stratification and processing of laser data are done, it’s possible to now do the regression analysis. There are a variety of statistical software programs that can be used to perform calculations, but for the results to be good requires both statistical and expertise. A regression function should be developed for each forest variable to be estimated, possibly divided per stratum. A first step is to decide what type of model is to be constructed. Residuals of regression should be evenly distributed. If the residuals are seen to increase or decrease for the independent variable, the data may be linearized, for example by using logarithms.

When selecting the independent variables, you can either start with one and gradually add new ones, or start with a lot of them and take away those which are not of importance. The former method is more manageable, then you should aim for a function with relatively few independent variables. One must also consider which laser measurements are reasonably associated with the variables to be estimated. For example, it is likely that wood volume is associated with both height and density measurements. Automatic methods such as stepwise regression and best subset regression, can provide guidance in the selection of variables. However, one should not rely blindly on the results because the methods are purely based on statistical relationships and do not necessarily say anything about the actual causation. The input laser measurements should not be too highly correlated with each other, such as two adjacent height percentiles. For each variable that is added or removed, a new, temporary, regression function is made. By analyzing the correlation between the residuals from the temporary regression function and each candidate for the new regression, it is determined which variable is to be added next. Table 9.2 shows examples of the regression functions which have been used in different studies to estimate different forest variables.
Table 9.2. Examples of regression functions for estimating the population level, from studies of Næsset et al. and Holmgren et al.

<table>
<thead>
<tr>
<th>Study</th>
<th>Variable</th>
<th>Funktion³</th>
</tr>
</thead>
<tbody>
<tr>
<td>Næsset</td>
<td>Mean height, H</td>
<td>( \ln(H) = 0.35 + 0.529 \ln(h_{90f}) + 0.355 \ln(h_{max}) )</td>
</tr>
<tr>
<td>Holmgren</td>
<td>Mean height, H</td>
<td>( H = 1.46 + 0.95 \ h_{95} )</td>
</tr>
<tr>
<td>Næsset</td>
<td>Stem volume, V</td>
<td>( \ln(V) = 3.151 + 3.027 \ln(h_{80l}) - 1.66 \ln(h_{max}) + 1.223 \ln(d_{50f}) )</td>
</tr>
<tr>
<td>Holmgren</td>
<td>Stem volume, V</td>
<td>( \ln(V) = -2.50 + 0.87 \ln(D_v) + 1.49 \ln(h_{90}) - 2.44 \text{relustd} + 0.44 \ D_p )</td>
</tr>
<tr>
<td>Næsset</td>
<td>Mean diameter, Dg</td>
<td>( \ln(d_g) = 0.406 + 0.892 \ln(h_{80}) - 0.374 \ln(d_{1f}) )</td>
</tr>
<tr>
<td>Næsset</td>
<td>Stem number, N</td>
<td>( \ln(N) = 10.33 - 0.487 \ln(h_{90}) - 0.667 \ln(h_{cvf}) + 1.187 \ln(d_{3f}) )</td>
</tr>
</tbody>
</table>

³) Subskript f and l indicate that the metric is calculated for only the first (f) and last (l) returns. \( h_{max} \) is the maximum height of vegetation hits, \( d_{50f} \) is the number of first returns from the treetops of \( H_{50} \) divided by the total number of returns. \( D_v \) is the number of first returns above 3 m divided by the total number of returns. \( \text{relustd} \) is the standard deviation in height divided by \( H_{95} \). \( D_p = (n1 + n3) / (n1 + n2) \), where \( n1 \) is the number one returns, \( n2 \) is the number of first returns, and \( n3 \) is the number of first returns where others return is over 3 m.

The complete regression functions (one for each estimated variable and strata) is applied to the grid of laser data for estimation of forest variables. The resulting screens in combination with stands limits in vector format used to calculate population averages. Results from some Swedish experiments with regression method are shown in Table 9.4.

So-called cross-validation is usually often used to evaluate the results. Cross validation will be more thoroughly addressed in Chapter 10 on Accuracy Assessment. But as an example “Leave-one-out-Cross-validation” works by leaving out one plot at a time (or sometimes many), estimates the functions parameters and uses these to estimate a value for the single plot or group of plots taken away. This is done for all plots and the estimation result is then compared with the plot field measurements.

Regression provides unbiased estimates, i.e., the value of the estimated variables on average are correct. Since the regression method is model-based and allows interpolation, regression works with relatively little field data. Sample plots must, however, be fairly representative - you should not, for example have too many plots that are unusually dense for their height.
Table 9.3. Results from some Swedish experiments with regression method, evaluated at the stand level and compensated for sampling error.

<table>
<thead>
<tr>
<th>Estimated variable</th>
<th>Mean Error (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Volume/ha</td>
<td>6-14</td>
</tr>
<tr>
<td>Basal-area weighted mean diameter</td>
<td>7-13</td>
</tr>
<tr>
<td>Stem number/ha</td>
<td>12-24</td>
</tr>
<tr>
<td>Basal-area weighted mean height</td>
<td>3-6</td>
</tr>
</tbody>
</table>

9.6.2. k-MSN and k-NN

The k-MSN (k Most Similar Neighbors) method is based on imputation. The k-MSN, and other similar methods such as k-NN (k Nearest Neighbors), selects a number of plots which have similar spectral values as the raster cell to be estimated and calculates a weighted average of these. The $k$ stands for the number of plots to use, and the value of $k$ can vary depending on the purpose, but typical values are between 1 and 20. A high $k$ gives better average estimation results, but may give less realistic relationships between the estimated variables in a given area.

The difference between kMSN and kNN is that kMSN uses canonical correlation analysis to select the neighbors. This part is a parametric method, which is based on the linear combination of the dependent variables (here forest data) that are most correlated with a combination of independent data (e.g., laser measurement) is calculated. This gives the weights for the different laser metrics importance in imputation.

All dependent variables are estimated while allowing the natural relationships between the estimated variables retained better than the regression. The estimation of k-MSN is widely used for estimations using remote sensing data from spectral data as well as laser scanning.

It is important to have a large number of test surfaces evenly spread over the entire range of variation. The greater the variation in the forest, the more plots are needed. Extrapolation works poorly because a theoretical model is not calculated in this case, and only the actual data in the data set are used to assign values. To illustrate, consider how kMSN or kNN might work for estimating wood volume over a large area if the training data only has values up to 200 m$^3$/ha.

The k-MSN method requires significantly more field data, evenly spread over the entire range of variation in the data. Since the method is based on imputation to each grid cell of forest data from plots that have similar characteristics as the laser data, it is very important that the extremes of the data set are represented.
9.7. Combining estimation and classification

Note that output from continuous value estimation can also be used to assign a thematic class to a pixel. As a simple example, a raster output of percent forest cover can be converted to a two class thematic map of forest and non-forest by defining forest using a threshold in the percent of forest cover (e.g., > 10%). When continuous values are estimated with the goal of creating a thematic map, this is sometimes referred to as “soft” classification, whereas a thematic classification is referred to as a “hard” classification (Fernandes et al., 2004).

9.8. Selecting the remote sensing data variables

As you know by now, satellite images have several wavelength bands of data. Which bands do you want to use in the classification process? This is also a subject of research. The answer is to use a combination of statistical analysis and common sense to decide which bands to use. Are there certain wavelengths or variables from the remote sensing data that are highly correlated with the vegetation property you wish to map? Then include this band. Are there several bands of remote sensing data which are highly correlated? Then you may want to use only one of these. Correlation analysis, best subset regression, variable importance or step-wise addition of information are just some of the methods used to choose the remote sensing data which should be used in the analysis.

We can look at the data visually. In a very simplified example, we can easily plot the values of the pixels in a 2-D graph, or scatter plot (Fig 9.8).

![Simplified two-band scatter plot showing the training data for different thematic classes.](image)

Figure 9.8. A simplified two-band scatter plot showing the training data for different thematic classes.

To look at more than two dimensions, we can look at the spectral signature of different land cover types plotted for all spectral bands (Fig. 9.9)
Figure 9.9. Spectral signatures of different land cover classes from the 15-band MERIS data.
9.9. Change analysis

Change detection is one of the large operational applications of optical satellite remote sensing of forests. The Swedish forest administration for example acquires medium resolution satellite data (SPOT) annually for all of Sweden. The main reason for this is that the images are used for checking the location and year for new clear felled areas. The areas where changes are detected are then matched with the cutting permits.

In other countries, such as Estonia, change detection has also been used for checking how large proportions of the fellings that have been made without cutting permits. Many countries are also discussing the use of change detection techniques for the mandatory deforestation mapping to be made for Carbon reporting to the UNFCCC.

Change detection is also an important technique for damage mapping. In Sweden, it has for example been used for mapping the extent of Gremeniella damages and for mapping areas affected by storm damage (for example, after Gudrun in southern Sweden).

To date, most change detection studies have been performed with aerial photos, optical satellite data, and radar data. As the use of laser data is relatively new and few multitemporal data sets exist, change detection studies with laser data has not yet been extensively used operationally.

9.9.1. Prerequisites for change detection

The comparison of imagery from different time points is a powerful tool for detection of local changes in the forest, compared to the normal development. Before such change detection can be carried out, a number of prerequisites, listed below, should be considered.

- Time of year, and number of years between images
- Pixel size
- Geometry
- Sensor type and spectral bands

Time of year, and time between images

For Swedish conditions, it is preferable that both images are from the mid- or late part of the summer, when the vegetation greenness is stable. Images from spring, early summer, or autumn, might give variations in color that depend on the tree species and these variations might be mistaken for changes. In practice, this means from about June 15 to August 31 for Sweden.

An additional factor is considering the phenomena you are observing, and selecting a number of years between images that makes sense. For example, if you wish to detect clearcuts in a tropical area, where forest regrows much faster than in a boreal forest, you will need to have images with less time between the acquisition dates.
Pixel size
In order to have as few “mixed pixels” as possible and to have a “many pixels per stand” situation, a small pixel size is preferable; on the other hand, “many trees per pixel” will give easily handled mean values. Thus, for the Swedish forest landscape, pixel sizes in the order of 20 m, and down to as small as approximately 5 m, is preferable.

Geometry
The images should be registered to the same co-ordinate system with the same pixel size, e.g. 20 * 20 m pixels, oriented squarely using the Swedish coordinate system.

Sensor type and spectral bands
It is preferable that the same sensor type and spectral band, is used at each time point, but that is not absolutely necessary; while most changes in the Swedish forest will be best visible using a shortwave infrared band, it might also be of some help to make change imagery from more than one spectral band.

9.9.2. Simple change detection techniques
The basic idea in most change detection applications within forestry is that within specific areas (often stands) changes can be seen when comparing earlier and later images. Some different techniques can be used, such as

- Visual display
- Post-classification change
- Image differencing.

Visual display
Visual display involves quickly switching between satellite images from two subsequent summers (e.g. images from July 2015 and July 2016), new clear-felled areas will appear to be brighter in the later image, whereas the rest of the forest landscape will appear quite similar in both images.

Another simple technique is to display data from different time points using different colour display guns in an image processing system. For example, one band from the July 2015 image is displayed on both the blue and green display colour gun and one of the corresponding bands from the July 2016 image is displayed using the red display colour gun. In this way, areas that have been brighter in the later image due to clear-felling, for example) will appear as red, while all other areas will have a rather common grey tone (note that colour balancing of the images is often needed to achieve the best result here).

Post classification change
This method consists of using classifications from two different images and judging the difference between the classes. This method may be seen in the literature, but is not the most accurate way to do change detection. The reason why it isn’t the most accurate is because each image will have classification
errors. When using both images together, the errors will be compounded, resulting in even more error in the result.

**Image differencing**

Major changes in the forest, like clear-fellings, will show up using very simple techniques listed previously. For detection of more marginal changes in the forest, a change image might be useful, this is what is commonly meant with “change detection” in remote sensing.

If we let

- \( T_1(j) \) = the pixel value for pixel \( j \) in the early image (image \( T_1 \))
- \( T_2(j) \) = the pixel value for pixel \( j \) in the later image (image \( T_2 \))

A usual way for computing a change image is then:

\[
\Delta_j = a \left( T_2(j) - f(T_1(j)) \right) + b
\]

Where \( \Delta_j \) = the pixel value for pixel \( j \) in the change image, \( f \) is a function that adjusts the pixel values in image \( T_1 \) to the same grey scale as the pixel values in image \( T_2 \), and \( a \) and \( b \) are just coefficients for contrast enhancement of the change image. Often \( b \) is chosen to be equal to 127.

The next question is then, which method should be used for estimating the calibration function (\( f \)). The reflectance calibration approach would be to transfer each image to the reflectance scale, and then work with the differences between reflectance calibrated images. However, this far, this method has not been accurate enough for forestry. Furthermore, there are also natural reasons (like seasonal differences) for reflectance differences between images. The method currently recommended is therefore to use a statistically based “relative” calibration, based on the grey values in each of the two images.

**9.5.3. Some statistical approaches for relative calibration of images to each other**

The next question is how to compute the function \( f \), that adjust the radiometry (= the pixel values) of the early image \( T_1 \), to that of the later image \( T_2 \). There are two main approaches: pixelwise methods and distribution based methods. Both methods are statistical and we can extract a number of pixel values from each image and do the computations on tabulated data, as illustrated below:

<table>
<thead>
<tr>
<th>Pixel</th>
<th>( T_1 )</th>
<th>( T_2 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>23</td>
<td>25</td>
</tr>
<tr>
<td>2</td>
<td>22</td>
<td>26</td>
</tr>
<tr>
<td>3</td>
<td>12</td>
<td>14</td>
</tr>
<tr>
<td>etc</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

In pixelwise methods, we use the pair-wise pixel values from \( T_1 \) and \( T_2 \) in an estimation method, most typically regression, where we give the pixel values for \( T_2 \) as the dependent variable, and the pixel values of \( T_1 \) as the
independent variable, thereby getting a regression function. The residuals between expected and actually measured values will have the same scaling as T2 and could be used as a change image. Advantages with regression analysis are that it is a flexible and easy understood standard method, and several bands in the early image could also be used to predict one single band in the later image. However, all pixelwise methods are sensitive to geometric errors between the two images T1 and T2. (One possible way to reduce this problem would be to work with segment mean values instead).

In distribution based methods, we only work with the summary statistics of the pixel values in each image, thus, these methods are not sensitive to geometric errors between the images. One such method, that works well and is implemented in most image processing systems, is histogram matching. The steps are to:

- compute separate cumulative histograms of both T1 and T2 with pixel values on the X-axis and the percent (%) of pixels within the image on the Y-axis;
- for a DN level in T1, read which % of the cumulative distribution it corresponds to;
- read which DN in T2 has the same cumulative %;
- make a look-up table with T1 DN values on the X-axis, and the values in T2 that has the same cumulative distribution value, on the Y-axis.
- repeat the three previous steps for all DN (0-255)
- update T1 through the look-up table to obtain T1’, which is an image with same grey value distribution as T2
- we can then compute a change image as
\[ \Delta j = a (T2_j - T1'_j) + b \]

The use of a forest mask

The next question is which pixel values to base the above procedures on? They will work for a sample of all pixel values in the image. However, for forestry change detection, they will (at least for Swedish conditions) work even better if they are based on a sample of forest pixels only, which can be defined by a digital map mask, for example.

9.5.4. Which forest changes will be visible in a change image?

The following image (Fig 12:35) comes from the textbook “Flygbildsteknik and Fjärranalys”. The image shows a change image made using two SPOT Panchromatic bands over Brännland, just NW Umeå. A 10 m pixel image from 1986 has been histogram matched against a similar image from 1989, and a difference image has been computed where the adjusted 1986 image has been subtracted from the 1989 image. The middle grey areas shows normally developed forest; the brighter areas are changes that usually are
associated with a biomass decrease, or exposure of mineral soil; the darker areas are associated with a biomass increase of a type that have a relatively large effect on the spectral signature.

The numbers in the image, are according to field checks, associated with the following changes:

1) the dark area is a young forest plantation with much deciduous trees, that during the 3 year period probably has passed the stage where it forms a closed canopy;
2) the bright area is a clearfelling;
3) the narrow white line is a newly dig ditch with exposed mineral soil, located on a clearfelled area;
4) the light grey area is a final felling in pine forest, with left seedling trees;
5) this light grey area is a final felling, where the seedling trees have been cut
6) this dark area was subject to final felling just before 1986, and since then, there have been a heavy increase in the grass cover;
7) this light grey area is young plantation where a deciduous shrub cleaning has been carried out between the image acquisitions.

In terms of reflectance factors (R), the following is typical for boreal forest in the Umeå area (Table 9.4):

<table>
<thead>
<tr>
<th>Table 9.4. Reflectance Factors from Landsat TM data.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
</tr>
<tr>
<td>Medium aged, pine dominated forest</td>
</tr>
<tr>
<td>Increase caused by strong thinning</td>
</tr>
<tr>
<td>Increase because of seed tree stands</td>
</tr>
<tr>
<td>Increase because of clear felled areas</td>
</tr>
</tbody>
</table>

In other words, the increase in reflectance due to clearfelling is about 10 times greater than that of thinning cuttings. Still, thinning cuttings are often visible in change imagery, especially in pine forest. However, from a change image only, it might be difficult to tell what the type of change it is.

Damages, for example from the fungi Gremeniella that damage pines, snow break damages, or wind thrown trees, can also be detected in change imagery. Under favorable circumstances (good summer images, and a very carefully made analysis) it might be possible to detect damages where only 20% of the basal area has been damaged. The spectral change of the above-mentioned changes are similar to those caused by thinning cuttings.

The detection of the existence of a change is quite easily done by interpretation of a change image. The delineation of the changed areas are also quite feasible. It is possible to use automated image processing techniques, like segmentation, or interactive line following. A quite crude procedure that often is used in practice, is interactive thresholding of the pixel values in the change image. It should however be observed that there are mixed pixels at the borders of changed objects. Furthermore, the forest surrounding a clear-felled area is sunlit on one side and causes shadows on the other side of the area, and therefore thresholding techniques may not necessarily result in correct delineation of clear-cut areas.

The labeling of the type of change is the most difficult part. Many changes will cause a similar spectral response, for example thinning cuttings, snow break damages, and gremmeniella fungi damages. The combined use of the change image and the early image, will contribute with some information about the type of forest before the damage. However, we have to accept that
it is much easier to detect the existence of change than to be able to automatically assign the reason for the change. In reality, field visits or more detailed information (detailed field inventories) are required for certain knowledge of the reasons for change. Automatic procedures will require field references, or calibrated images and known thresholds.

9.10. **Time series analysis**

Another way to use remote sensing data from different time points is to use data from several different time points to answer questions. For instance, by using a number of Landsat images over several decades (e.g., imagine a series of images from 1975, 1980, 1985, 1991, 1996, 2002, 2008, 2012, 2017), we may be able to distinguish the relative age of a forest after a clearcut, or can observe the development of the forest by looking at trajectories (creating a vector of values by observing a single pixel over many time points). We can expect in this type of application to observe forest that has grown in height and density at different rates over the landscape, and can then label them as productive or non-productive forest classes. In another application, 3D remote sensing data can be used to look at forest height development over time, and thereby say something about the forest productivity.

Another way to use a time series is to use satellite images from many dates over different seasons, and in this way, capture phenological changes of the vegetation that help to identify that class more accurately. An example would be the use of this tactic to identify different types of deciduous tree species that have different bud-break and senescence times (e.g., oak and alder).

9.11. **Data fusion**

There are two ways to look at the term “data fusion”. One is quite literally, and an example of this is the process of pan-sharpening. Pan-sharpening is a technique used to combine spatial higher resolution bands with coarser spatial resolution bands. For example, Landsat 7 and 8 have a panchromatic band with a 15 m pixel resolution, while the multi-spectral bands have a 30 m pixel resolution. By using a weighting technique, the information in the four 15 m pixels contained in the multi-spectral 30 x 30 m pixel can be fused to create a multi-spectral dataset that has the appearance of a higher spatial resolution.

Another way to consider “data fusion” is more the approach of using multiple sources of data in a classification or estimation. This might be by using both Landsat-8 and Sentinel-2 data, as well as a DEM. This isn’t a “true” data fusion, as you could refer to this simply as using multiple data sources, but it may sometimes be referred to in the literature as data fusion.
9.12. Data assimilation

9.13. Software used for image processing
Currently there is a movement from use of commercial packages towards open source code for processing of remotely sensed data. For optical satellite data, some common commercial packages are Erdas Imagine; ENVI; IDRISI; and eCognition (especially for segmentation). ArcMap is primarily a GIS software, but due to earlier collaboration with Erdas it has some limited image processing capabilities. QGIS is also increasing its capabilities to include image processing routines. Several commercial software have the ability to write and implement code, such as ENVI’s IDL interface. The more commonly available open source software are provided through R and OSGeo4W, although the number of these available are foreseen to increase over the next decade. Finally, those with a knowledge of a programming language, such as C++, can also write their own code to use for image processing.

Statistically, common classification methods, (for example maximum likelihood classification, which as you read earlier is actually a form of Discriminant Analysis) can be found in statistical packages (like SAS, MINITAB, SPSS etc), instead of an image processing system. While this isn’t necessary, because most image processing systems (like ERDAS Imagine) have classification routines, it is possible. These days it is becoming more common to use the statistical program R for image classification activities.

Självstudiefrågor

Litteraturen
10. ACCURACY ASSESSMENT

10.1. Accuracy assessment of remote sensing data products

Accuracy assessment of the map product is often an important element for the users of the data. It is sometimes not carried out, however, for large area projects, due to the lack of reference data, or limitations in project time or funding. An independent and objectively collected evaluation data set is essential to an unbiased assessment of the map product. Accuracy assessment numbers give the user insight into the average accuracy of the map, the accuracy of different classes/categories, and why errors may be happening.

10.1.1. The need for probability sampling

Data for accuracy assessment should be collected with a method that is much more accurate than the product to be evaluated. In this case field visits (with accurate GPS positions), or interpretation of aerial photos are most often used for accuracy assessment.

Accuracy assessments should be done with a sample that is objectively selected according to the rules of probability sampling. This means that a random sample of evaluation points is perfectly valid, and also a systematic sample is quite OK from most aspects. It is also allowed to use a stratified approach, where a different number of evaluation points is used for different strata. However, what is not allowed in probability sampling, is to sample reference plots in a subjective way, such as “where it is convenient”, or to “move” plots that happen to be located in less favorable places. If there is a need to avoid certain types of plot locations, rules for this have to be created in advance (for example after a pilot survey), and the accuracy assessment figures are then only valid with the restrictions given by those rules. An important characteristic of validation data is that it should have high quality measurements (position and inventory measurement).

10.2. Assessing thematic class accuracy

The number of accuracy assessment plots needed is often an issue, and the answer to that question often needs a statistical analysis that is not part of this course. However, there is a rule of thumb, where 50 samples per class are recommended (Congalton 1991).

10.2.1. The error matrix

The most usual way to analyze and present an accuracy evaluation is to construct an error matrix, also called a confusion matrix or contingency
The error matrix is a table with a row for each class in the classification, and a column for each category according to the field (or photo) evaluation. A simple example of an error matrix is given below:

<table>
<thead>
<tr>
<th>Data according to classification:</th>
<th>Field data</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Coniferous forest</td>
</tr>
<tr>
<td>Coniferous forest</td>
<td>30</td>
</tr>
<tr>
<td>Deciduous forest</td>
<td>3</td>
</tr>
<tr>
<td>Non-forest</td>
<td>4</td>
</tr>
<tr>
<td>Total</td>
<td>37</td>
</tr>
</tbody>
</table>

The “producer’s accuracy” = how large a proportion of the field data for a given class was correctly classified:
- Coniferous forest: \( \frac{30}{37} = 81\% \)
- Deciduous forest: \( \frac{10}{17} = 59\% \)
- Non-forest: \( \frac{30}{52} = 58\% \)

The “user’s accuracy” = how large a proportion of the classified data for a given class had the correct label according to the field data:
- Coniferous forest: \( \frac{30}{52} = 58\% \)
- Deciduous forest: \( \frac{10}{15} = 67\% \)
- Non-forest: \( \frac{30}{39} = 77\% \)

The ratio between Producer’s and User’s Accuracy for a single class can reveal whether a class has over-classified or under-classified, according to the reference data. For example, there were many more instances of coniferous forest in the classification than in the reference data. Coniferous forest is “over-classified”.

The most used measure of classification accuracy is overall accuracy, which means the total number of correctly classified pixels, divided with the total number of pixels, in the above table, the overall accuracy is estimated as:

\[
\frac{30 + 10 + 30}{106} = 66\%
\]

In general, an overall accuracy of 80% for about 8 land cover classes is rather reasonable.

The Kappa coefficient (\( \hat{k} \)) or k-hat is a measure where the overall accuracy have been reduced with the accuracy that only depends on the chance agreement between the classification and the field evaluation. The measure has a value between 0 and 1, and is always lower than the overall accuracy. The detailed formula is given in Lillesand et al. The general formula is:
k-hat = \frac{(\text{observed accuracy} - \text{chance agreement})}{(1 - \text{chance agreement})}

10.3. Assessing continuous estimate accuracy
For continuous estimates RMSE is a common way to assess error.

The RMSE can be calculated as:

\[ RMSE = \sqrt{\frac{\sum_{i=1}^{n} (\Delta_i)^2}{n}} \]

where \( \Delta_i = (\hat{v}_i - v_i) \)

\( \Delta_i \) is differences between estimated parameter \( \hat{v}_i \) and the field measurements (or “truth”) of the same parameter \( v_i \) for the \( i \)th observation (in our previous lab each observation was each stand in the database), and \( n \) is the number of observations.

Relative RMSE is comparing the error of the estimated parameter as a percent to the mean value of the field measured parameter. For example, if the RMSE of the estimate was 10 and the field measured parameter’s mean was 100, then the relative RMSE would be 10%.

Bias can also be calculated as \( \frac{\sum (\hat{v}_i - \bar{v})}{n} \)

10.4. Cross-validation
Cross-validation is a method used when independent data are not available to conduct a separate accuracy assessment, because the data have been used for training or model creation. Leave-one-out cross-validation removes one of the plots from the reference data set, creates the model with the remaining data, and assesses the accuracy for the single plot not used. The procedure then goes through the entire data set in this manner, and the results are compiled. Cross-validation gives one a good idea of model-performance, and is not a “true” assessment of the result, but can be used as an indication when there is a limited reference data set to work with.

10.5. Considerations about the collection of field data
There will be geometry errors between the field data and the image data, which will reduce the apparent classification accuracy if a “pixel wise” evaluation is done. Furthermore, it is often difficult to obtain a correct class label for just a point in field, at least without a lot of time consuming measurements. To judge a class label for a bit larger area is often easier. A classified map itself is also more accurate if a mean value for a segment is
used. These are some arguments about why it might be better to evaluate accuracy for groups of pixels (e.g. segments with the size of stands) instead of single pixels. Furthermore, a judgment in field might often be near the class definition between two different classes. There are also ways to compute an error matrix where classes that are very similar can be considered less wrong than classes that are very different, which is called “fuzzy accuracy assessment”.

Other tricky topics are how to get accuracy assessment plots for change detection? Some possible data sources are serial photos from past time points, long term inventory series, or you might simulate a change.

### 10.6. Further reading

Stehman and Czaplewski (2003) defined four criteria that should be met: 1) probability sampling, 2) adequate sample sizes with which to estimate user’s accuracies with acceptable level of precision, 3) cost efficiency must be considered, and 4) spatial distribution of samples must be representative across the area of interest.

Stehman and Czaplewski (1998) have established three basic elements to consider in the design of an accuracy assessment plan: the sampling design, the response design, and the estimation and analysis protocol. The sampling unit may be a pixel, fixed-area plot or polygon, although the optimal unit depends on the application. Stehman *et al.* (2000) favor pixel-based evaluation units, as larger units render the results non-site specific. Polygon assessments also tend to lead to conservative estimates of classification accuracy (Verbyla & Hammond, 1995). Class homogeneity within the accuracy assessment unit is appealing, but not necessary, and if intentionally included, may bias the assessment of the map accuracy. A design-based sample with known inclusion properties is best, but the distances between plots should be large enough that potential spatial autocorrelation effects do not influence the result. Definitions constituting correct and incorrect responses should be established (e.g., if polygon accuracy assessment units are used, the rule may be that a “correct” classification requires a majority of the classified pixels to be correctly labeled as the dominant class, or the rule may be that the two most dominant classes must both be classified).

As with training data, questions regarding the sample size and sampling scheme of the accuracy assessment data need to be addressed. Stehman (2001) suggests that a sample size of 100 samples per class assures the population is estimated adequately. Congalton and Green (2009) suggest a minimum of 50 samples per class. To capture the necessary number of samples for rare classes, a stratified sample may be useful (Stehman, 2001).

The quality of evaluation data is of much importance, and a quality check of the evaluation data should be carried out before use.
Accuracy assessment is often presented in an “error matrix”, with errors of commission (called “user’s accuracy”) and omission (called “producer’s accuracy”) for each class in the map, as well as a measure of overall accuracy. The kappa statistic, or k-hat, is also a measure of overall accuracy, and is intended to account for the chance of random agreement (Congalton and Green, 2009). A desired overall map accuracy of 85% is often given as a benchmark, but may not be realistic to achieve (Wulder et al., 2006). Fuzzy accuracy assessment (Gopal & Woodcock, 1994; Foody, 2002) can be a useful measure of portraying different types of errors that may be more or less acceptable.

It is slightly more common that inventory data are used for accuracy assessment than for training (e.g., Riemann et al., 2010). Wulder et al. (2006) encountered difficulties when applying 2 ha polygon-based inventory data due to differences between the raster and vector data, particularly because the polygon interpretation included heterogeneous cover. When purpose-collected video data were later photo-interpreted for accuracy assessment, the uncertainty in the photo-interpretation and the lack of a probability-based sample were drawbacks (Wulder et al., 2007). One of the primary requests emerging after Canada’s EOSD land cover mapping project was for improved collection strategies of calibration (training) and validation (accuracy assessment) data (Wulder et al., 2008).
11. TILLÄMPNINGAR AV FJÄRRANALYS

11.1. Globala karteringar

11.2. Nationella skogskartor

Satellitbildsskattningar med stöd av rikstaxytor
I de Nordiska länderna har flera Rikstäckande skogliga karteringar gjorts genom sambearbetning av satellitbilder från Landsat och liknande satelliter, och med användning av provytor från Rikskogstaxeringar i respektive land. Denna utveckling startade med etableringen av Finlands ”Multisource National Forest Inventory” i början på 1990-talet (Tomppo et al., 2008). Med den så kallade kNN metod som då tillämpades, så tilldelades varje pixel alla skogliga data från den eller de provytor som hade mest lika färg som provytan. Resultatet blir en rasterdatabas med skogliga variabler för alla rasterrutor inom skogsmark. När data från flera rutor aggregeras, så kan statistik beräknas för mindre områden än när bara rikstaxrutor används. Denna ”multi source” metod används operativt i den Finska Riksskogstaxeringen för beräkning av skoglig statistik på kommunnivå.

I Sverige gjorde SLU år 2000 en liknande produkt som i Finland, med Landsat ETM+ satellitbilder och rikstaxytor som tilldelades satellitbildspixlarna med kNN metoden (Reese et al., 2003). SLU har sedan upprepat denna kartering år 2005 och 2010, med data från SPOT satelliterna. Dessa data är fritt tillgängliga från Riksskogstaxeringens hemsida, de kallas officiellt ”SLU Skogsskarta”, men går också under namnet kNN-Sverige.

Även i Norge har Riksskogstaxeringen gjort en liknande landstäckande satellitbildsprodukt kallad SAT-SKOG (Gjertsen, 2007). Det har också gjorts liknande produkter i Danmark.

Denna typ av landstäckande skogsdatabaser har använts för en mängd ändamål, t.ex. som datakälla för ekologisk forskning och som översikt för myndigheter. Det förefaller alltid att finnas ett behov av den för tillfället bästa kompletta kartan av ett lands skogsresurser. För operativt skogsbruk, så har dock satellitbildsskattningar som baserar sig på data från enstaka tidpunkter inte ansetts bra nog. Därför är produkter som bygger på 3D punktmoln av större intresse, se nästa avsnitt.

Landstäckande skogliga skattningar med stöd av 3D data från laserskanning eller digital fotogrammetri
Precis som det går att göra skogliga skattningar i nationell skala med rikstaxdata och satellitbilder, så kan 3D punktmoln från nationell
Laserskanning användas istället för satellitdata. Skattningarna av variabler som medellödhöjd, medeldiameter, grundytta och virkesförråd blir då väsentligt bättre än om endast ”2D” satellitdata av Landsat-typ används.


**Skogliga Grunddata från Skogsstyrelsen och SLU**

Lantmäteriet påbörjade år 2009 en rikstäckande laserskanning av landet med syfte att ta fram en ny nationell höjdmodell. Skanningen har utförts på ca 2000 m flyghöjd och punkttätheten i markplan är 0.5 – 1 punkter/m². Vid skanningen har Sverige delats in i 387 block, vanligen med en storlek om 25 x 50 km (Figur 1). Ambitionen har varit att varje block ska skannas inom en kort tidsperiod med samma skanner. Totalt har 13 olika skannerindivider använts för den nationella skanningen. De flesta blocken har skannats med Leica- (74 %) eller Optech-skannerar (25 %) och endast ett med skannerar från Trimble- eller Riegl. Tanken har varit att skanna södra Sverige, som har mer lövträd, på våren och hösten när löven fallit, medan norra Sverige mestadels skannats under sommarhalvåret. Anledningen till denna uppdelning är att täta trädkronor i lövskogar kan hindra lasern att träffa markytan.

Laserpunktmolnen ger inte bara information om marken utan även om vegetationens höjd och täthet. Detta har uppmärksammats av Regeringen som gett Skogsstyrelsen i uppdrag att i samarbete med SLU och i dialog med andra intressenter ta fram rikstäckande skogliga skattningar utifrån Lantmäteriets laserdata.

Under perioden 2013-2016 har rikstäckande skattningar av virkesvolym, grundytta, medellödhöjd, medeldiameter och trädbiomassa ovan mark tagits fram genom att kombinera Lantmäteriets laserdata med provytedata från Riksskogstaxeringen. Detta har gjorts med hjälp av regressionsmodeller som tagits fram genom att först beräknas statistiska mätt (metriker) för hur laserträffarna inom Riksskogstaxeringens provytor fördelas sig i trädskiktet.
och på marken. Därefter har regressionsmodeller som skattar de aktuella
variablerna eller transformationer av variablerna utifrån lasermetrikerna
tagits fram. Dessa har sedan använts för att skatta variablerna för varje
rasterruta i landskapet om 12,5 x 12,5 meter, där även de aktuella
lasermetrikerna beräknats. Skattningarna har tagits fram för ett block i taget
 genom att först skatta om regressionsparametrarna för modellerna som
äterges i tabell 1. Därefter har den modell för respektive variabel som gav
högst förklaringsgrad (R²) för respektive variabel använts. Eftersom antalet
provytor inom ett block normalt är relativt litet användes även provytor
med tillhörande laserdata från närliggande block som skannats med samma
skannerfabrikat och under samma skanningssäsong (dvs. med eller utan löv
på träden). De framtagna uppgifterna redovisas endast för rasterceller som
har en skattad grundytevägd medelhöjd om minst 3 meter och för block
som huvudsakligen ligger nedanför Skogsstyrelsens gräns för
föryngringsavverkning.

Skogliga grunddata består av rasterkartor med 12,5x12,5 meter stora
rasterceller där det för varje rastercell finns skattade värden för volym,
grundyta, trädbiomassa, grundytevägd medelhöjd och grundytevägd
medeldiameter. Dessa data kan laddas ned fritt från Skogsstyrelsens
hemsida. Resultatet kan sedan aggregeras till större områden, t.ex. bestånd,
genom att man beräknar medelvärdet av alla rasterceller som ingår i ett
bestånd. Varje användare kan själv rita ut de bestånd som ska analyseras,
t.ex. med de verktyg för detta som finns på Skogsstyrelsens hemsida.

I tillägg till de skattningar för 12,5x12,5 m som rutor som gjorts av SLU, så
har Skogsstyrelsen även låtit ta fram ytterligare produkter från den
nationella laserskanningen och som finns på deras hemsida. Exempel är:
trädhöjdsraster för 2x2 m rasterceller, markfuktighetskarta,
terrängskuggning och slutningskarta. En del av dessa produkter säljs sedan
tidigare även av konsulter och har därför av marknadsskäl inte gjorts fritt
dedladdningsbara, åtminstone inte inledningsvis.

Tabell 1. Regressionsmodeller som använts i projektet skogliga grunddata

<table>
<thead>
<tr>
<th>Variabel</th>
<th>Regressionsmodell*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gy-vägd medelhöjd (m)</td>
<td>( h_{bw_i} = \beta_0 + \beta_1 h_{p95} + \varepsilon_i )</td>
</tr>
<tr>
<td>Stamdiameter (cm)</td>
<td>( d_{bw_i} = \beta_0 + \beta_1 h_{p80} + \beta_2 (h_{p80} * vr) + \varepsilon_i )</td>
</tr>
<tr>
<td></td>
<td>( d_{bw_i} = \beta_0 + \beta_1 h_{p90} + \beta_2 (h_{p90} * vr) + \varepsilon_i )</td>
</tr>
<tr>
<td></td>
<td>( d_{bw_i} = \beta_0 + \beta_1 h_{p80} + \beta_2 (h_{p90} * vr) + \varepsilon_i )</td>
</tr>
<tr>
<td>Grundyta (m²/ha)</td>
<td>( b_{ai} = \beta_0 + \beta_1 (h_{p80} * vr) + \beta_2 \text{Std}(lh) + \varepsilon_i )</td>
</tr>
<tr>
<td></td>
<td>( b_{ai} = \beta_0 + \beta_1 (h_{p90} * vr) + \beta_2 \text{Std}(lh) + \varepsilon_i )</td>
</tr>
<tr>
<td>Stamvolym (m³/ha)</td>
<td>( \sqrt{v_i} = \beta_0 + \beta_1 h_{p80} + \beta_2 (h_{p80} * vr) + \beta_3 \text{Std}(lh) + \varepsilon_i )</td>
</tr>
<tr>
<td></td>
<td>( \sqrt{v_i} = \beta_0 + \beta_1 h_{p90} + \beta_2 (h_{p90} * vr) + \beta_3 \text{Std}(lh) + \varepsilon_i )</td>
</tr>
<tr>
<td></td>
<td>( \sqrt{v_i} = \beta_0 + \beta_1 h_{p80} + \beta_2 (h_{p90} * vr) + \beta_3 \text{Std}(lh) + \varepsilon_i )</td>
</tr>
<tr>
<td>Biomassa (ton/ha)</td>
<td>( \sqrt{b_{ioi}} = \beta_0 + \beta_1 h_{p80} + \beta_2 vr + \beta_3 \text{Std}(lh) + \varepsilon_i )</td>
</tr>
<tr>
<td></td>
<td>( \sqrt{b_{ioi}} = \beta_0 + \beta_1 h_{p90} + \beta_2 vr + \beta_3 \text{Std}(lh) + \varepsilon_i )</td>
</tr>
<tr>
<td></td>
<td>( \sqrt{b_{ioi}} = \beta_0 + \beta_1 h_{p80} + \beta_2 vr + \beta_3 \text{Std}(lh) + \varepsilon_i )</td>
</tr>
<tr>
<td></td>
<td>( \sqrt{b_{ioi}} = \beta_0 + \beta_1 h_{p90} + \beta_2 vr + \beta_3 \text{Std}(lh) + \varepsilon_i )</td>
</tr>
</tbody>
</table>

*) \( h_{px} = \) höjdpercentil x; \( vr = \) antal förstarturer över 1.5m / (totalt antal förstarturer).
Figur 2 visar resultat av validering av skogliga grunddata på beståndsnivå. Som framgår av tabellen, så är noggrannheten för skattning av volym strax under 20%. Resultaten kan därmed anses vara i nivå med, eller något bättre, än de manuella skattningar som görs i samband med skogsbruksplanläggning. Dessa goda resultat har också bidragit till att produkten blivit accepterad i skogsbruket och ofta används som ett stöd vid skogsbruksplanläggning. Den används även av många andra aktörer, t.ex. banker och försäkringsbolag.

Skogliga grunddata möjliggör smidig och kostnadsfri beräkning av skogliga data för valfritt område. Det finns dock en rad begränsningar, bland dessa märks:
- trädslag och ståndortsindex är inte skattade eftersom det inte kan beräknas från den typ av data som använts;
- ingen avgränsning av den produktiva skogsmarken har gjorts, utan den måste användaren själv definiera;
- de skogliga uppgifterna avser respektive områdes skanningsår, nu har dock Skogsstyrelsen utvecklat ett verktyg som möjliggör en enkel framskrivning från skanningsåret;
- bestånd med mycket höga förråd kan vara underskattade och bestånd med låga förråd kan vara överskattade, så kallad ”dragning mot mitten” som förekommer vid alla liknade skattingsmetoder;
- bestånd som skannats under lövad säsong kan vara överskattade om de innehåller mycket lövträd.

Genomförandet av projektet har inneburit att ny teknik har utvecklats för storskalig och effektiv produktion av skogliga data. Två andra statliga satsningar, Lantmäteriets laserskanning för en ny höjdmall samt Riksskogstaxeringens provytor har genom projektet fått ett stort mervärde. Fritt tillgängliga Skogliga grunddata, som med god noggrannhet både beskriver skog och mark, är starten på någonting nytt som ger många positiva följeeffekter, som ökad effektivitet och bättre kvalité i skoglig planering. Utvecklingen mot nya användningsområden, exempelvis för att hitta områden med höga naturvärd, har också tagit fart.

### 11.3. Datafångst för skogsbruksplanering

#### Manuell datafångst för skogsbruksplanering

Den viktigaste fjärranalysdatakällan vid skogsinventering i Sverige är fortfarande flygbilder, i synnerhet som stora delar av Sverige numera flygfotograferas varannat år. Som beskrivs närmare i kapitel 5, så kan flygbilder antingen användas för förhandstolkning i fotogrammetrisk arbetsstation eller, vilket är mycket vanligare idag, i form av ett digitalt ortofoto.

Vid manuell tolkning av ett digitalt ortofoto mot bildskärm eller läsplatta, så är det också enkelt att växla till andra dataskikt. Exempel på nya typer av dataskikt från laserskannerdata som visat sig användbara för skogsbruksplanering är terrängskuggning, som bl.a. kan visa diken och kulturminnen; markfuktighetskartor, samt högupplöst trädhöjdsraster som ger god information om beståndsgränser. Av dessa så kan trädhöjdsrasteret även framställas från digital fotogrametri.

Skogliga data samlas oftast in i fält med hjälp av relaskop och höjdmätare, men även om indelningen och mätning görs manuellt så kan givetvis automatiskt beräknade skattningar som skogliga grunddata ändå utgöra ett stöd vid skogsbruksplaneringen.

#### Laserskattningar för skogsbruksplanering

Den operationella användningen av laserdata för skoglig planering startade i Norge år 2002 (Naesset et al, 2004). Där hade man redan ett system för att samordna skogsbruksplaneringen kommunvis. Detta system består och för närvarande betalar staten 57% av kostnaden för planerna, vilket gör att de flesta markägare ansluter sig till den gemensamma planläggningen när det är aktuellt att göra planer i deras område. Planerna utförs av konsultföretag och arbetet görs med ett minimum av fältbesök. Före laserskanningens genombrott fångades så mycket som mycket som möjligt av datat genom tolkning och mätning i fotogrammetrisk arbetsstation, dvs det som vi i Sverige kallar ”LMV-metoden”. Indelning i avdelningar och uppskattning av trädslag görs fortfarande manuellt med fotogrammetrisk arbetsstation, medan de flesta övriga variabler fångas från flygburna laserdata. Vid denna typ av projekt brukar särskilda provytor mätas strax efter skanningen. Denna typ av inventeringar kan därför anpassas bättre till laserskattningarnas behov. Provytorna kan t.ex. läggas där de gör mest nytta, t.ex givet att man delat in skogen i ett antal strata, med avseende på trädslag och stådortsindex och vill ha ett visst antal provytor per strata. Denna typ av specialanpassade laserskanningsprojekt bör sammantaget ge något bättre skattningsnoggrannheter än om rikstaxdata används som referens, hur mycket bättre det blir och om det är värt de extra kostnaderna är dock en öppen fråga i dagsläget.


11.4. Habitatkartering


Mänskliga aktiviteter har förändrat ekosystemen i hela världen, vilket har lett till habitatförlust och nedgång för många arter. Modern skogsbruk med homogena bestånd av träd med samma art och ålder har ersatt stora områden med naturligt föryngrad skog. Detta har lett till fragmentering av habitat och minkande habitatkvalitet för många arter. För att planera och sköta skog för att gynna biologisk mångfald krävs kunskap om olika arters habitatkrav.

Information som kan härledas från fjärranalysdata är användbar även för att beskriva habitat. För habitat i skogsmiljöer är bland annat trädslagssammansättning, struktur och ålder relevanta. Vegetationsstruktur är en egenskap som har betydelse för biologisk mångfald och som kan vara enklare att beskriva från fjärranalysdata än från fältinventeringar. Fjärranalys kan användas för att få information om förhållandena på en plats men även för att beskriva fördelnings av områden med vissa egenskaper i landskapet. Detta kan användas för att skapa kartor över potentiella habitat för olika arter. En annan användning är att skaffa kunskap om vilka habitatkrav en art har genom att analysera i vilka miljöer...

**Laserdata för habitatkartering**


Fjärranalys beskriver typiskt sett egenskaper hos vegetationen med relativt grov upplösning, till exempel trädslag eller höjd- och diameterfördelning för träden. Detta är viktigt information, men även andra faktorer har betydelse, till exempel artsammansättning hos övrig flora och fauna eller brukningshistoriken (Brumelis et al. 2011). Detta är svårare och ibland omöjligt att beskriva med fjärranalys. Ett annat problem är att måået kan vara att modellera lämpliga habitat, men eftersom de exakta egenskaperna hos lämpliga habitat inte är helt kända används istället ofta observationer av arterna. Risken med detta är att utbredningen av arterna inte motsvarar utbredningen av lämpliga habitat (Bradley et al. 2012). Det kan finnas lämpliga habitat dit arten inte har kunnat eller hunnit sprida sig och det kan finnas områden som inte är optimala habitat dit arten har spridit sig från lämpliga habitat i omgivningen.

**Referenser till kapitel 11.2 och 11.3**


forest variables using satellite data and field data from the National Forest Inventory. *Ambio* 32: 542-548.


**Referenser till kapitel 11.4**


Appendix

Internet Länkar

Allmäna länkar
- Rymdstyrelsen - http://www.snsb.se/
- European Space Agency - http://www.esa.int/ESA
- CEOS - http://ceos.org/
- Indian Space Research Organization - http://www.isro.gov.in/
- Google Earth Engine - https://earthengine.google.com/#intro

Remote sensing and forests
- Natural Resources Canada - https://www.nrcan.gc.ca/forests/measuring-reporting/remote-sensing/13429
- Global Forest Watch - http://www.globalforestwatch.org/

Data Access
- SACCESS – https://saccess.lantmateriet.se/
- Sentinels Science Hub - https://scihub.copernicus.eu/
- Lantmäteriet - http://www.lantmateriet.se/
- Global Land Cover Facility - http://www.landcover.org/

Tutorials
- Natural Resources Canada - http://www.nrcan.gc.ca/earth-sciences/geomatics/satellite-imagery-air-photos/satellite-imagery-products/educational-resources/

Organizations
- EaRSEL - http://www.earsel.org/
- ISPRS - http://www.isprs.org/
- ASPRS - http://www.asprs.org/
Sweden specific websites
- Kiruna Ground Station - http://www.esa.int/Our_Activities/Operations/Estrack/Kiruna_station
- Swedish Space Corporation and ESRANGE - http://www.sscspace.com/

Journals and newsletters
- Space News - http://spacenews.com/

Miscellaneous
- One hundred applications of remote sensing - http://gisgeography.com/100-earth-remote-sensing-applications-uses/
- TED Talk, Greg Asner “Ecology from the air” - https://www.youtube.com/watch?v=qCrVpRBBSvY
- Ian Woodhouse, “What is remote sensing?” - https://www.youtube.com/watch?v=8HhfJsiYenE
- How does LiDAR work? - https://www.youtube.com/watch?v=EYbhNSUnIdU

Our own websites
- Ljungbergsfond - http://ljungbergsfonden.se/
- Section of Remote Sensing at SLU – www.slu.se/srh
- The Ljungbergs Lab at SLU – www.rslab.se

Kapitel 1: Introduktion till skoglig fjärranalys

Kapitel 2:

The Electromagnetic Spectrum
- https://www.esa.int/SPECIALS/Eduspace_EN/SEM7IQ3Z2OF_0.html
- http://www.seos-project.eu/modules/remotesensing/remotesensing-c01-p01.html

Color additive theory