Andrei-Cristian Baraian

PARTICLE SIZE DISTRIBUTION BASED ON DEEP LEARNING INSTANCE SEGMENTATION

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ABSTRACT

Deep learning has become one of the most important topics in Computer Science, and recently it proved to deliver outstanding performances in the field of Computer Vision, ranging from image classification and object detection to instance segmentation and panoptic segmentation. However, most of these results were obtained on large, publicly available datasets, that exhibit a low level of scene complexity. Less is known about applying deep neural networks to images acquired in industrial settings, where data is available in limited amounts. Moreover, comparing an image-based measurement boosted by deep learning to an established reference method can pave the way towards a shift in industrial measurements.

This thesis hypothesizes that the particle size distribution can be estimated by employing a deep neural network to segment the particles of interest. The analysis was performed on two deep neural networks, comparing the results of the instance segmentation and the resulted size distributions. First, the data was manually labelled by selecting apatite and phlogopite particles, formulating the problem as a two-class instance segmentation task. Next, models were trained based on the two architectures and then used for predicting instances of particles on previously unseen images. Ultimately, accumulating the sizes of the predicted particles would result in a particle size distribution for a given dataset.

The final results validated the hypothesis to some extent and showed that tackling difficult and complex challenges in the industry by leveraging state-of-the-art deep learning neural networks leads to promising results. The system was able to correctly identify most of the particles, even in challenging situations. The resulted particle size distribution was also compared to a reference measurement obtained by the laser diffraction method, but still further research and experiments are required in order to properly compare the two methods. The two evaluated architectures yielded great results, with relatively small amounts of annotated data.

Keywords: neural networks, non-intrusive, minerals, machine learning
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FOREWORD

This master’s thesis was written at VTT Technical Research Center of Finland, Oulu, while I was part of the Machine Vision team. I would like to express my gratitude to Professor Janne Heikkilä from University of Oulu and Vili Kellokumpu from VTT for supervision and guidance. I would also like to thank Outotec for providing valuable feedback and providing data for this work. I am grateful for everyone at VTT for the support and ideas during this whole project. Also, I would like to thank Janne Mustaniemi for providing the second supervision. Finally, I would like to thank my family and friends for their unconditional support and encouragement.

Oulu, March 10th, 2021

Andrei-Cristian Baraian
### LIST OF ABBREVIATIONS AND SYMBOLS

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>AI</td>
<td>artificial intelligence</td>
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<tr>
<td>ANN</td>
<td>artificial neural network</td>
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<tr>
<td>CMOS</td>
<td>complementary metal–oxide–semiconductor</td>
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<tr>
<td>CNN</td>
<td>convolutional neural network</td>
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<td>DNN</td>
<td>deep neural network</td>
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<td>DL</td>
<td>deep learning</td>
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<tr>
<td>FPN</td>
<td>feature pyramid network</td>
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<td>FPS</td>
<td>frames per second</td>
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<td>GAN</td>
<td>generative adversarial network</td>
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<td>IoU</td>
<td>intersection over union</td>
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<td>KDE</td>
<td>kernel density estimation</td>
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<td>mAP</td>
<td>mean average precision</td>
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<td>ML</td>
<td>machine learning</td>
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<td>MLP</td>
<td>multi layer perceptron</td>
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<td>NN</td>
<td>neural network</td>
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<td>NMS</td>
<td>non maximum suppression</td>
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<td>PSD</td>
<td>particle size distribution</td>
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<tr>
<td>ReLU</td>
<td>rectified linear unit</td>
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<td>RNN</td>
<td>recurrent neural network</td>
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<td>ROI</td>
<td>region of interest</td>
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<td>RPN</td>
<td>region proposal network</td>
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<tr>
<td>SGD</td>
<td>stochastic gradient descent</td>
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<td>SNR</td>
<td>signal to noise ratio</td>
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<td>SVM</td>
<td>support vector machine</td>
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<td>UE</td>
<td>ultimate erosion</td>
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1. INTRODUCTION

Particle size distribution (PSD) estimation is a well-known technique applied in many industrial sectors for monitoring, controlling and optimization of various essential processes. In the mining industry, the grinding process is responsible for reducing the particle size by a combination of impact and abrasion, in dry environment or more commonly, in suspension of fluid. It is also the last stage of the comminution process, which is a mechanical process of size reduction for solid materials [1, 2, 3]. To monitor and control the grinding circuit reliably, the size distribution of particles needs to be constantly estimated, preferably in real-time.

The first method developed for the task of particle size distribution estimation was sieve analysis, where particles of interest are run through a stack of sieves having different size dimensions. The PSD is computed by weighting the amount of material stopped by each sieve. A simple and efficient method, it provided decent enough results for a long period of time. However, with the proliferation of non-intrusive methods based on optical devices, more complex and robust solutions have been designed. By using a laser beam oriented towards the particle, the laser diffraction method can be applied, which relies on the angle and intensity of the scattered light to estimate the size of a particle. Especially for small particles, ranging from a few nanometers to millimeters, it proves to be a very efficient and accurate method. Digital image processing represents another emerging technique used to estimate the size distribution of particles by leveraging the recent developments in machine vision cameras. In this case, the particles are directly identified in images and their real-world size is obtained by converting the pixel size into a metric size. Moreover, Deep Learning (DL) architectures are becoming increasingly popular even in industrial computer vision applications, boosting traditional image processing techniques to achieve better performance.

In this thesis, the focus is centered on image-based analysis methods for PSD estimation applied in mining scenarios, which use state-of-the-art deep learning architectures for instance segmentation. The general pipeline for obtaining the PSD consists of individually identifying the particles of interest in the image, thus obtaining a mask for each particle. This process is known as instance segmentation. After obtaining the masks, the area for each particle is computed and together with the camera parameters and the employed camera model, its real-world size can be estimated. The instance segmentation module is usually implemented with traditional image processing techniques, but lately, DL has been successfully applied for this task and achieved better results than any other method. Although the performance of DL models in solving computer vision tasks is well-documented [4], less is known about their performance when applied on complex data collected in industrial settings and how well they satisfy the requirements of industrial applications. Primarily, this study evaluates the capability of a Deep Neural Network (DNN) to efficiently segment the instances of mineral particles, with the final purpose of estimating the particle size distribution. The main advantages of using a DNN rather than a traditional image processing pipeline or machine learning (ML) approach are the reduced set of parameters that need to be configured, increased accuracy, robustness and speed. The only parameters that need to be adjusted are the network parameters, but they are independent of the data on which it is trained, hence allowing for easy transfer to
new data. Furthermore, rather than designing complex flows for corner-cases, we only have to make sure that the training data consists of few corner cases, so the focus is shifted from algorithm development to data analysis, which in most cases can be more accessible.

Continuous monitoring and control tasks usually need to adhere to specific time, latency and accuracy requirements. Since there are multiple DNN architectures that solve the problem of instance segmentation, but have certain key differences, two of them will be compared in-depth, taking into consideration the requirements of the presented use case. Therefore, the main contributions of this thesis are:

- A pipeline for obtaining the size distribution of particles from mining images, namely apatite and phlogopite, using a DNN for generating the instance segmentation.

- Comparison of two DNN architectures considering the task of instance segmentation and the resulted particle size distribution.

In the next section, the thesis introduces the context of particle size analysis in the mining sector and gives a brief overview of the most popular methods for particle analysis, with a strong emphasis on image-based methods. Section 3 starts describing the underlying foundations of DL and presents the two DNNs architectures used in the thesis. Then, previous work and results related to DL and particle size analysis are discussed. Section 4 highlights the implemented pipeline and the data analysis, while Section 5 shows the obtained results and both the comparison of the two architectures and the comparison of the DL methods against the laser diffraction method. The last two sections are critically analyzing the obtained results, further research directions are proposed and the whole work is summarised.
2. PARTICLE SIZE ANALYSIS - OVERVIEW

Mining has been performed for thousands of years across the whole world, and it is the backbone of sustaining and developing the infrastructure of our societies. With the huge increase in raw materials demand for emerging technologies and infrastructures, it becomes clear that we need more efficient mineral excavation and processing, such that we have a sustainable framework in which to operate, now and in the future.

Extracting useful minerals is a demanding and complex process. Most of the time, valuable minerals are mixed with other non-valuable or useless materials and for separating them, first we need tools to distinguish them. A particle analyzer is one such tool that can analyze and report information about the size distribution of particles in a sample. The results are then used for subsequent control and monitoring of various mining processes such as grinding. Choosing the right particle analyzer depends on some key parameters such as: size ranges, chemistry/material of the particles, desired information, performance requirements. Of course, there are other indirect parameters that can influence the decision, like budget, current analysis technique, etc.

There are three main types of particle analyzers, each relying on different technology and having their own advantages and disadvantages. The most rudimentary technique is sieve analysis, which works for reasonably sized particles, and it is a mechanical, intrusive process. If we want to continuously analyze particles, especially small ones, then we have to use either laser diffraction or image-analysis based methods. This chapter will present an overview of these methods, highlighting the way in which they are able to calculate the particle size distribution and the environment in which they operate.

2.1. Sieve Analysis

Sieves have been used for a very long time in the mining industry. It provides a quick and easy way of measuring the particle size for a large number of particles, instead of individually picking particles and having a human operator measuring them. This was the first step towards automating the process of measuring particles. The system consists of a stack of sieve meshes placed vertically, like a column. The mesh (screen) at the top has the largest screen openings and subsequently the lower levels have smaller screen openings than the one above. The stack of meshes is placed inside a shaker, which shakes the structure for a period of time. Then, on each mesh, the weight of the material is measured and combining the results, a PSD is obtained.

This method is one of the most used one, mainly because of its simplicity, efficiency and low cost. Also, the technology has evolved and there are sieve analyzers that perform quite well in terms of accuracy, reliability and processing time. But, it is still a mechanical process and particles can be affected by the impact with the mesh grid, leading to some particles breaking in smaller pieces, therefore affecting the size distribution. Moreover, the sieve meshes can suffer as well from the impact leading to some screen openings getting larger and not preserving the size consistency for a particular level in the mesh stack. From the measurement point of view, it lacks in precision, since it assumes rectangular shaped objects and most of the time grinding particles have irregular shapes. Furthermore, there are cases when elongated objects
can fit through a small opening if they happen to be oriented in a specific way. It is not an online measurement, since we need to perform the shaking for some time, wait for the particles to settle and only after that we can get the results. Another drawback is the limit on particle size that can be measured. Really small particles cannot be measured since they are too small for this mechanical process, and it is impossible to design such fined-grained sieve meshes which operate in the range of nanometers.

2.2. Laser Diffraction Analysis

![Laser diffraction particle measurement. Reprinted with permission from Outotec.](image)

When particles are getting exceptionally small, somewhere in the range of a few submicrons to millimeters, laser diffraction technique is the way to go. The working principle behind this concept is depicted in Fig. 1. The particle flow is placed between a laser light source and a detector. The laser beam is diffracted by the particles at different angles, depending on the particle’s size and the scattered light is focused by a lens on a concentric array of photodetectors. The particle size is obtained by measuring the angular variation in intensity of the light scattered on the detector. Larger particles scatter light at a lower angle relative to the laser beam than smaller particles. The scattering pattern is then interpreted for getting the actual size of the particle using either Mie [5] or Fraunhofer theory.

It is clear that such a procedure is much more complex than sieve analysis, due to the increased cost of operation that it brings. It allows for faster and continuous measurements, high throughput, increased accuracy, etc. Like sieve analysis, laser diffraction also expects particles to have a pre-defined shape, which is spherical in this case. Moreover, laser diffraction is a non-contact method, hence the particles maintain their physical structure.
A cutting-edge particle size analyzer that uses laser diffraction is Outotec’s PSI 500 Particle Analyzer. It provides real-time PSD estimation for particles in slurry environment, usually used for monitoring grinding circuits, regrinding circuits, backfill and tailings disposal and feed to the slurry machine. Fig. 2 shows the physical build of the PSI 500 device along with a sample PSD computed by the system.

![PSI 500 Particle Size Analyzer](https://www.mogroup.com/portfolio/psi-500i-particle-size-analyzer/?r=2)

Figure 2. PSI 500 particle size analyzer device and an estimated PSD. Reprinted with permission from Outotec [6].

### 2.3. Image-Based Analysis

With the proliferation of machine vision algorithms and recent advances in cameras, image analyzers have gained attention from the mining industry as well. Acquiring images in an industrial environment is considered to be a challenging task due to harsh conditions such as sudden temperature changes, dust or vibrations. New imaging technologies are aiming at mitigating the artifacts introduced by these external factors, and image-based analysis promises to be a very efficient measurement tool. The stakeholders involved in mineral processes may be tempted to use machine vision solutions for control processes due to being inexpensive, fast, non-intrusive, consistent, robust and accurate.

The general pipeline of an image-based analysis starts by acquiring the raw image with a camera sensor and the necessary illumination setup, such that the particles of interest can be clearly distinguished in the image. The second step and usually the most complex one consists in segmenting the image, obtaining a binary image that can discriminate between objects (in our case, particles) of interest and background or other non-related objects. After obtaining the segmented image, we can calculate different size measures for each particle and obtain a size distribution across a batch of samples, images in our case, so that the number of particles is statistically sufficient.

There are four main problems in computer vision related to identifying objects of interest from images. A visual understanding of these problems is depicted in Fig. 3.

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Figure 3. Different aspects of object identification in computer vision.
(1) Image classification (2) Semantic Segmentation (3) Object localization (4) Instance Segmentation

Image classification was the first problem proposed, and it is simply aimed at solving the task of identifying the class of the main object in the image, regardless of its position, orientation or other objects. There is an assumption that only one class is assigned for an image, representing the main object. Object localization can identify and localize the objects at the same time, enclosing them in bounding boxes. It makes the transition from image-level classification to instance-level classification. Semantic segmentation operates in a slightly different way and performs the classification at pixel level. As in the figure above, we are interested in segmenting the rocks from all the other objects and background. But semantic segmentation does not tell us how many objects there are in the picture, although this can be solved by additional post-processing algorithms. The last problem, instance segmentation is one of the most difficult one, and it builds on top of object localization, adding a segmentation mask for each detected instance. Depending on the employed technique for size measurement and on the object of interest to be detected, each technique has its own advantage. The one having the most information is clearly instance segmentation, but when comparing it with object localization, most networks designed for instance segmentation are slower since it adds the overhead of calculating a segmentation mask for each detection.

To get an overview of the segmentation and localization methods used for estimating the PSD, we can categorize them as follows: conventional, ML based and Convolutional Neural Network (CNN) based methods [7].

Conventional methods rely on traditional image processing techniques for creating a pipeline that can segment the particles of interest. Popular methods include Watershed Transform [8], ultimate erosion (UE) [9] and the Hough Transform [10]. The first two methods can prove to be quite fast and utilize low memory, but are more prone to errors
and susceptible to noise in images. On the other hand, Hough Transform can be more robust, but is slower and has a large memory footprint. The big disadvantage shared by all these conventional methods is having to adjust parameters by the user, depending on the imaging conditions, the particles to be observed and other major changes in the operating environment. These type of algorithms are unfortunately not robust enough by themselves and any change can potentially lead to erroneous results, meaning that the system needs to be recalibrated each time. Moreover, fine-tuning the parameters can be time-consuming and if performed frequently, it can become a major bottleneck.

**ML-based methods** [11, 12] are one step closer towards making the process more autonomous, since there are very few image or environment dependent parameters that need to be set by the user. ML-based methods rely on two key concepts, namely feature extractors and classifiers. The classifiers are trained with the extracted features, finding patterns in the input data. Then, based on the learned features, the classifiers will label unseen data. Descriptors are usually used as input data to the ML classifiers, meaning that a lot of effort is shifted towards designing efficient and robust descriptors, capable of capturing as much information as needed. By compressing parts of images in descriptors, the available information is reduced, affecting the accuracy of the model.

**CNN-based methods** are end-to-end methods that can learn not only feature interpretation, but feature extraction as well. This means that there are virtually no parameters that need to be set by the user, as CNN methods work directly on the raw image. On the other hand, they are heavily dependent on vast amounts of annotated data, so that the model can extract useful information. Getting enough data for every application where CNNs are used can be very tricky and most of the time even impossible. But, a good technique to overpass the lack of data is transfer learning, where a model is trained on general purpose data with the objective of learning the feature extraction and later training only the last layers on the specific, reduced dataset for our application. This will be discussed in detail later on.

There are also hybrid methods, that combine the principle of laser diffraction with imaging techniques. In [13], the authors built a particle analyzer using a CMOS image sensor and a collimated beam configuration, together with a ML model based on Random Forest. Similarly to laser diffraction, they have used the angular distribution of scattered light to measure particle size.

### 2.4. Limitations of Traditional Image Processing Methods

Image-based analysis proves to offer enough information and data for estimating the PSD accurately and robustly, but even after choosing this approach, there are a wide range of algorithms that can achieve the segmentation of an image. Conventional methods based on traditional image processing techniques were the founding blocks of image-based analysis. But, they soon proved to be difficult to develop, maintain and transfer the same concept or pipeline to a new setup, even though the requirements were the same. One of their main disadvantages is the huge number of parameters involved. Each step in a general pipeline requires several parameters to be adjusted, for example, image smoothing, edge detection, thresholding, morphological operations, labelling, etc. Not only does this make the development hard, but a slight change in the working environment may affect the pipeline altogether and crash the system.
Moreover, none of these methods can compete with the accuracy of a human operator [11].

A more robust approach is to use machine learning algorithms, that are able to better predict properties of objects in the image or even classify them. These algorithms have an increased tolerance for changes in the working environment and can better handle unknown data. But, ML algorithms are also dependent on the extracted features and part of the workload does not disappear, but is actually shifted towards designing robust and efficient feature extractors and descriptors. While having far fewer parameters to tune than traditional image processing techniques, there is still a number of parameters that need to be set for the feature extractors and classification models and sometimes designing the perfect features extractor can be quite time consuming.

Because of all these inconveniences, end-to-end methods are really desired in the industry. Working conditions can change, especially in industrial environments, so increased flexibility is needed. Also, there are a lot of mineral processes that have a high degree of similarity and rather than designing a completely new system for each in turn, it would make sense to transfer the core concept and adjust as few parameters as possible. DNNs are pushing forward the state-of-the-art in computer vision and are enabling end-to-end methods, where users only need to feed annotated data and then the network is capable, on its own, of extracting the features, learning them and ultimately classifying new data.
3. DEEP LEARNING BASED INSTANCE SEGMENTATION

The computer vision community has seen a great increase in the utilization of DL in last years and starting from the success of AlexNet [14] in the ImageNet competition (Large Scale Visual Recognition Challenge, ILSVRC 2012 [15]) it has since been the hot-topic of computer vision. Not only has DL extended to other tasks in computer vision such as object detection and localization, semantic segmentation, instance segmentation, pose estimation, and so on, but it has been successfully applied in other domains as well, such as audio [16], natural language processing (NLP) [17], 3D reconstruction [18] and many others. Keeping in mind the recent advancements in terms of GPU processing capabilities, it becomes more and more clear that DL is for now, the tool of choice for solving the problems of today and tomorrow.

Because DL is such a vast domain, the next section will be focused on a brief overview of the core idea behind DNNs and how they have evolved in the context of machine learning and under the big umbrella of artificial intelligence. The explanations will start from the building blocks of DNNs, then gradually going through state-of-the-art architectures. Popular DL models will be presented, with applicability in the instance segmentation domain.

3.1. AI Vs. ML Vs. DL

The concept of Artificial Intelligence was born in the 1956 by the AI pioneers who envisioned complex machines capable of expressing human intelligence. This means sensing, interacting with the real-world and taking human-like decisions. Conceptually, this is known as ’General AI’ and would ideally be represented as machines that can behave and reason like humans do. For now, this theoretically and maybe frightening concept cannot be achieved, but instead, more specialized forms of AI have been developed for particular scenarios and tasks, like image classification, text recognition, natural language processing, robot control, etc. These are still tasks or actions that require human-like intelligence, hence the algorithms based on DL are built specifically for a task, rather than building a ’general AI’ robot that can solve everything.

Machine learning is a subset of AI practices that shifts the paradigm from hand-programming to data learning. Rather than designing algorithms to behave in a certain
way, ML algorithms learn from the available data and then make educated guesses on unseen data. This consists of parsing the data, learning different patterns that are likely to occur by extracting features from data and then make a prediction about previously unseen data. ML encapsulates a wide set of algorithmic approaches, like clustering, reinforcement learning, decision-tree learning, feature-based learning, etc. Although it is more robust and flexible than previous methods, it still involves a lot of hand-coding and crafting of feature extractors.

Parts of the ML tool set are also Artificial Neural Networks (ANNs). They are inspired by the way in which our brain functions, leveraging the inter-connection between multiple neurons. In a Neural Network (NN), the neurons have weights associated with each of them and represent how wrong or correct it is relative to the performed task. Even though they were present from the early days of AI, the hardware was not yet ready for supporting this concept. The most basic networks were too computationally intensive and they were simply not feasible. With the deployment of GPUs, specialized hardware for parallel processing, the breakthrough of ANNs and specifically very deep ANNs (DNNs) was possible. Networks with increased number of layers and neurons were possible to train in reasonable times and some architectures proved to perform even better than humans in some tasks (image classification). But there is a twist. Deep neural networks require huge amounts of data, usually being trained on hundreds of thousands of images which requires them to be labelled. As the task increases in complexity, e.g. from image classification and up to instance segmentation and panoptic segmentation, the labelling process becomes more difficult and requires more time and resources. Also, when applying DL to a specialized, narrow domain, like mineral detection, there are few challenges like manual annotation, lack of available data, complexity of labelling, ambiguity in labelling and others.

3.2. Neural Networks

The building block of a neural network is the neuron. Inspired from biology, the neuron takes as input stimuli from multiple connections and when sufficient stimuli, it fires on the output. Likewise, in ANNs, a node (neuron) multiplies input data with a set of weights associated with every connection, trying to amplify or dampen that input, based on its significance relative to the performed task. This allows the network to characterize which input is helpful when classifying. The sum of product between weights and data is passed through an activation function that decides if or how much of the signal should pass onward. This can be mathematically formulated as:

\[
y = \theta(b + \sum_{i=1}^{N} x_i w_i)
\]  (1)

where \(y\) is the output of the activation function, \(x_i\) are the inputs, \(w_i\) are the weights of the networks, \(b\) is the bias and ultimately, \(\theta\) is the activation function. Visually, a node representation is depicted in Fig. 5a.
A NN is formed by stacking layers of neurons and connecting the layers. A generic architecture for a NN consists of an input layer, which consumes the input data, hidden layers and ultimately an output layer responsible for the prediction (represented as probability). A DNN is basically a NN that has many hidden layers, hence the name 'deep'. The motivation behind building deep neural networks is that the deeper you go in a neural net, the more complex features nodes can learn, since they are aggregated with previous nodes. Training is an iterative process that updates the weights and biases such that the loss function is minimized and the data is learned.

### 3.2.1. Activation Functions

If we think about Eq. 1 and ignore the activation function, a neuron is just computing the weighted sum of its input, obtaining a real number. Same as a biological neuron, an artificial neuron needs to decide whether it fires or not (it is activated or not). The mechanism allowing this is the activation function. There are two types of activation functions: linear and non-linear functions. The non-linear functions are mostly used due to their ability to generalize or adapt to the variety in data. Below, some of the most used activation functions are presented, alongside their graphs.

The sigmoid function is defined by:

\[ f(x) = \frac{1}{1 + e^{-x}} \]

It outputs a value between 0 and 1, it is non-linear, continuously, differentiable and monotonic, all being desirable properties of an activation function. A big drawback is the insignificant change in gradient for inputs that are far from the origin. This gives rise to the problem of vanishing gradient, where the network is not capable of properly learning anymore or the training becomes increasingly slower for saturated neurons.

A similar function, the hyperbolic tangent, is also a non-linear function, but unlike the sigmoid, it is zero-centered, resulting in a better mapping of negative/positive values, making them strongly negative/positive. Unfortunately, it also shares the problem of vanishing gradient. Its mathematical definition is:
Rectified Linear Units or ReLU, may sound complicated, but it can be easily formulated as just:

\[ f(x) = \max(0, x) \]

It is non-linear, and has the same advantages as the sigmoid, but proves to have better performance. Also, it does not suffer from saturation and it is much easier and faster to compute the gradient. On the other hand, it is suffering from the well-known problem of "dying ReLU". In the case of having an input \( x < 0 \), the gradient will be 0, which means the weights will not be adjusted. Hence, those neurons will stop responding to variations in input. Leaky ReLU is a variant of ReLU where a small, non-zero gradient \( \alpha \) is allowed when the input is below zero.

![Graphs of activation functions.](image)

Figure 6. Graphs of activation functions.

3.3. Architecture of DNN

As it was mentioned in Section 3.1, general AI is rather a sci-fi dream than reality and the same is true for DL. There is no 'good-for-all' architecture and because of that, a range of different techniques have been developed for specific tasks that can boost the efficiency of a DNN. In this section, the most representative techniques are presented.
3.3.1. Convolutional Neural Networks

Convolutional neural networks (CNN) or ConvNets, have gained massive attention in computer vision and serve as basis for state-of-the-art architectures like AlexNet [14], VGGNet [19] or GoogleNet [20]. The components of a CNN are usually: convolution layer, pooling layer and fully connected layer. One of their main advantages is the capability of capturing both the spatial and temporal dependencies in an image by applying various filters. Since the core concept of ConvNets relies on reducing the number of parameters, therefore retaining only the most important features, CNNs can better fit the data than traditional NNs.

Convolutional layer

The convolutional layer is the most important part of a CNN. It relies on the convolution operation [21] to convolve the input image with different filters and obtain activation maps, which are in turn convolved again and again. Similarly, as in a neural network, a neighborhood or spatial region of the image, regarded as the input is convolved (multiplied) with the filter coefficients (weights). Then, an activation function is applied and the results are getting propagated through the network, similarly to a NN. The main objective of the convolution operation is to extract the high-level features such as edges or blobs. A convolution layer consists of a stack of filters, each with weights $w_i$, that will be the weights of the neural network that have to be learned. The spatial region on which we apply the convolution is called the receptive field. Different than in a Multi Layer Perceptron (MLP), where the output of a neuron depends on all the values from the previous layer, in a convolutional layer, the output depends on the filter $i$, specifically its weights, and the receptive field on which it is applied. The convolution operation usually decreases the dimensionality of the convolved feature, although padding can be used to either increase the dimensionality or keep it the same. In Fig. 7, a convolution with two 5x5x3 filters is applied.

Figure 7. Convolution of a 32x32x3 image with two 5x5x3 filters with stride 1 and no zero padding, that produces two activation maps.

The key parameters for a convolutional layer are the stride and padding. The stride is used for selecting the overlap between receptive fields and the padding, which is usually performed with zero values, is used to increase the size of the input, so pixels
situated near the border are considered as well. Using the following equation, one can determine the output of a convolutional layer:

\[(w + 2 \cdot p - k)/s + 1,\]

where \(w\) is the size of the input, \(k\) is the size of the kernel, \(p\) is the size of the padding and \(s\) is the stride.

**Pooling layer**

The pooling layer is responsible for down-sampling the input, reducing the spatial dimensionality. This is desirable for concentrating the information and retaining only what is considered to be most important, such that the network can learn more efficiently. Usually, it is placed after a convolutional layer. Also, by down-sampling, we obtain multi-resolution maps that can yield different important features. The most used operator is the max-pool one, retaining only the maximum value in the receptive field. Min and (global) average pooling are also applied, but are less frequent [22]. There are also studies in favour of dropping the pooling layer, especially when training Generative Adversarial Networks (GANs) or variational AEs, in favour of using convolution with bigger strides [23, 24].

![Figure 8. Pooling operation down-samples the input. Here, the input is downsampled using a max-pool filter with stride 2.](image)

**Fully connected layer**

The fully connected layer is usually used in the last stage of a CNN and is often responsible for outputting class membership probabilities. It works in the same way as a neuron in a MLP, taking into account all the input nodes from the previous layer. The last vector obtained by convolution and pooling is usually flattened and then fed to the fully connected layer, where a softmax function is applied.

### 3.3.2. Residual Block

A rule of thumb in deep learning was that the deeper the network is, the better it performs in terms of accuracy. This technique was preferred by researchers because of
its simplicity, but it proved to work only until a certain depth level. If the network is too deep, then problems like vanishing/exploding gradients or degrading appear [25, 26]. Vanishing gradient can happen if the weight initialization or data preprocessing steps are not done properly and also when applying the chain-rule over many stacked layers, then the gradient will get to zero eventually, resulting in the network not learning properly. Another problem that happens in Recurrent Neural Networks (RNNs), is the exploding gradient, where if we unroll the RNN for a number of steps and observe what the backward pass is doing, we can see that the gradient signal is getting multiplied many times with the same matrix, which can lead to the gradient ‘exploding’. Certain techniques have been developed to combat these problems and enable networks to converge using stochastic gradient descent (SGD) [27] by using normalized initialization and intermediate normalization layers [28] (batch normalization). Degradation, in terms of training accuracy, was observed when more layers were added, but the accuracy saturated or even started to decrease dramatically [29].

![Residual block](image)

Figure 9. Residual block. Copyright © 2016, Reprinted with permission from IEEE.

This problem was addressed by K. He et al. [30], which introduced a deep residual learning framework and were the winners of ILSVRC-2015 [15] classification task. Instead of simply stacking multiple layers on top of each other in the hope of learning an underlying mapping, they fit a residual mapping. This is done by adding a skip connection, also known as identity shortcut connection, as in Fig. 9. In their case, the skip connection just performs an identity mapping adding, where their outputs are added to the outputs of the convolutional layers. This way, instead of learning the desired underlying mapping \( H(x) \), the eq. \( F(x) + x \) is learned, where \( F(x) := H(x) - x \) is the residual mapping and authors claim that it is easier to learn the residual mapping than the original one. Hence, this framework enables the network to be trained end-to-end using backpropagation with SGD and no extra parameters or computational complexity is added by the shortcut connections. Moreover, the residual learning framework is generic and can be applied to other architectures as well.

3.3.3. Feature Pyramid Network

One of the biggest problems in the object detection task is being able to recognize objects at different scales. Especially for small objects, it can be hard for detectors to recognize them. A popular technique consists in building a pyramid of different image scales [31]. Then, processing each level in the image pyramid, objects at different scales can be detected. In practice, this approach has some drawbacks. Regarding performance, building an image pyramid takes significant time and memory and this
creates a problem for real-time applications. Also, as we go higher in the pyramid, the resolution decreases and the semantic value increases because of the high-level structures detected, but we cannot use bottom layers since the semantic value is low, although the resolution is higher.

A robust solution for this problem is the Feature Pyramid Network (FPN) [32]. FPN is a feature extractor that satisfies the requirements of speed and memory. As it can be seen in Fig. 10, it consists of a bottom-up and a top-down pathway, with lateral connections. The bottom-up pathway is pretty intuitive, enriching semantically each higher level while decreasing the resolution. The top-down pathway aims at reconstructing higher resolution layers from semantic rich layers. In terms of accuracy, the reconstructed layers are not so precise because of the upsample/downsample operations, so lateral connections are used to better predict the locations of objects. Similar as in ResNet, the lateral connections act as skip connections, making training easier. As a result, the architecture is able to build a feature pyramid with rich semantic features at all levels from a single input image scale. Another great advantage of FPN is being a generic solution that can be applied to many problems and architectures, like RPN (Region Proposal Network), R-CNNs (Region based Convolutional Neural Networks) or extracting masks for image segmentation.

![Figure 10. FPN Architecture. Copyright © 2017, Reprinted with permission from IEEE.](image)

### 3.4. Related Work

This section will give an overview of the evolution of Mask R-CNN, starting from the simple R-CNN and analyzing all the intermediate architectures which eventually led to the creation of the aforementioned. Yolact architecture will also be discussed as an alternative to Mask R-CNN and key differences will be highlighted, especially regarding those that affect the processing time.

#### 3.4.1. From R-CNN to Faster R-CNN

Object detection in computer vision is still one of the greatest challenge, since it is also the foundation of other tasks, like instance segmentation or size distribution of objects. Because we cannot assume the number of objects to be detected in an image, it is impossible to use a ConvNet with a fully connected layer and have as input only
one image. Multiple patches from an image could be used, but the question is how to choose them smartly in terms of aspect ratio and spatial location, since a brute-force method would be too computational intensive.

In [33], R. Girshick et al. proposed R-CNN, which stands for Regional-based Convolutional Neural Networks. R-CNN uses selective search [34] to extract only 2000 regions from an image which will represent region proposals. Their system consists of three modules. The first one is the region proposal generator, which fabricates 2000 candidate regions. Then, each region is fed to a CNN, which acts as a feature extractor and outputs a fixed-length vector as output. The last module is a set of class-specific linear SVMs, that classifies each region and also regresses four values, representing an offset of the bounding box, such that the location precision of the bounding box is increased.

**R-CNN: Regions with CNN features**

1. Input image
2. Extract region proposals (~2k)
3. Compute CNN features
4. Classify regions

The same author, in [35] managed to solve the problem of processing many region proposals by feeding the input image to the CNN, which generates a convolutional feature map. Then, region of proposals are identified from the convolutional feature map and a RoI pooling layer reshapes them into a fixed size by warping, so that they can be connected to a fully connected layer. Now, having a RoI vector, a softmax layer is introduced to predict the class and in parallel, a bounding box regressor finds the optimal offset. The new architecture is depicted in Fig. 12. These modifications allow the training to be single-stage, using a multi-task loss and the training can update all network layers.

Another notable attempt at speeding up R-CNN was the use of SPP nets [36]. Like Fast R-CNN, the SPP net computes a convolutional feature map from one image,
computes a feature vector for each object proposal and then classifies it using SVMs. Again, we have a multi-stage pipeline that takes significantly more time to train.

![Faster R-CNN Diagram](image)

Figure 13. Faster R-CNN. Copyright © 2017, Reprinted with permission from IEEE.

The authors of [37] proposed a new method for selecting the region proposals. Both R-CNN and Fast R-CNN were still using selective search, a bottleneck for the whole network because of the slow processing time. Therefore, after a convolutional feature map is obtained from the image, a separate network is used to learn the region proposals. The network is known as Region Proposal Network (RPN). Ultimately, proposals are fed into the RoI pooling layer for reshaping and are then classified and offsets are predicted. One of the main advantages of RPN is that it can be trained end-to-end by stochastic gradient descent. Starting from the foundation of R-CNN, Faster R-CNN became the distilled version which is significantly more accurate in predictions, but also achieves real-time inference performance. All these advancements paved the way for efficient instance segmentation, which is a much more complex task.

### 3.4.2. Mask R-CNN

After the huge success of Faster R-CNN architecture, it was clear that somewhere along those lines, future architectures for object detection and instance segmentation would need to be developed. And this was the case of Mask R-CNN [38], which is an extension of Faster R-CNN. With the addition of a mask head, it is capable of segmenting each detected object. So, instead of having just a bounding box around the object, the model computes a pixel-wise segmentation of that object. The novelties of Mask R-CNN are: the use of FPN [32], replacing ROIPool with ROIAlign layer and an additional branch that generates masks. A visual representation of the architecture is depicted in Fig. 14.
Figure 14. Mask R-CNN framework [38]. Copyright © 2017, Reprinted with permission from IEEE.

Mask R-CNN shares the same two-stage type procedure as Faster R-CNN. In the first one, RPN is responsible for generating region proposal, then in the second one, the class and bounding box offsets are predicted in parallel, with the addition of a mask predictor that works in parallel. Because of that, the multi-task loss is defined as:

$$L = L_{cls} + L_{box} + L_{mask}$$

The mask branch is unique in the sense that it generates masks for every class, hence there is no competition among classes and the classification is left to the class predictor. By doing this, mask and class predictions are decoupled, unlike usual FCNs [39], where a per-pixel softmax and multinomial cross entropy loss is used. Different than Fast R-CNN, pixel-to-pixel alignment is of high importance, which led the authors to propose ROIAlign, a layer responsible for aligning the extracted features with the input and getting rid of the aggressive quantization introduced by ROIPool.

A closer look at the architecture reveals that RPN is not applied on the original input image, but rather a backbone is used for extracting feature maps and subsequently, RPN scans over the backbone feature map. A backbone is a convolutional neural network that acts as a feature extractor, producing feature maps, which are passed further down the network. In terms of backbone architectures, the authors experiment with ResNet [30] and ResNeXt [40]. They also use a FPN [32], such that RoI features are extracted at different levels of the feature pyramid, resulting in substantial accuracy and speed improvement.

In [41], authors discover an inconsistency in assigning the score of an instance mask by using the box-level classification confidence. The confidence score takes into consideration only the difference between the semantic categories, and it is oblivious to the quality of the instance mask. For example, we might get a good bounding-box localization and a high classification score, but the mask quality can vary. This can further impact the evaluation and the training procedure. In the COCO [42] challenge, the evaluation is done by taking the average precision (AP) metric that uses Intersection-over-Union (IoU) score between the prediction and the ground-truth mask, but that is calculated for a fixed confidence threshold score. So, inspired by this, the authors propose a network capable of directly learning the mask IoU score and combining it with the confidence score as well, resulting in an alignment between the mask quality and its score. By addressing the problem of instance scoring, their solution improves the Mask R-CNN framework.
3.4.3. Yolact

Initial experiments using Mask R-CNN for the task of particle detection proved to be promising, but there was another requirement that had to be taken into consideration for the purpose of the overall system and that is the speed of processing. To efficiently monitor the PSD in the grinding process, real-time processing is mandatory, specifically in our case, we would need to process at least 10 frames per second (FPS). Bearing in mind this requirement, using Mask R-CNN is not feasible anymore since it can process 5 FPS on a beefy GPU. For this, the attention is shifted towards an architecture that is able to offer real-time processing without sacrificing too much accuracy. And Yolact [43] is the perfect candidate.

Fig. 15 depicts a comparison in terms of FPS vs. mAP on the COCO dataset. It can be easily seen that Mask R-CNN has a very good mAP score, but it lacks in speed. On the other hand, Yolact offers the best trade-off between accuracy and speed of processing. Another aspect that needs to be taken into consideration is the GPU on which the performance results are computed. All these comparisons have been done on beefy GPUs and considering our use case, it may be more feasible to have more compact processing units, like embedded GPUs, which do not have the same high processing power and memory capacity, so it is expected for the FPS rate to decrease when the model is deployed on such GPUs.

Previously, it was shown that Mask R-CNN is a two-stage detector, that relies on feature localization to generate masks. Only after the features have been re-pooled in the bounding-box region, they can be fed to the mask predictor, resulting in a sequential pipeline that represents a bottleneck in speed. To overcome this, D. Bolya et al. proposed Yolact [43], a single stage instance segmentation model capable of achieving real-time performance.

In order to achieve real-time performance, the authors break up the task of instance segmentation in two, parallel stages that are finally combined with minimum overhead. The overall architecture is presented in Fig. 16. The common parts of the network are
the backbone used for feature extraction and the FPN for producing more robust masks and high resolution prototypes. Then, in parallel, prototypes and mask coefficients are generated.

![Yolact Architecture](image)

Figure 16. Yolact Architecture [43]. Copyright © 2019, Reprinted with permission from IEEE.

Protonet is the network responsible for the generation of $k$ prototypes. It is implemented as a FCN which has $k$ channels in the last layer and uses as input a backbone feature layer, enhanced by the FPN. Although this looks similar to a semantic segmentation task, it differs by having no loss over the prototypes, instead the optimization is done from the final mask loss, that is calculated after assembly. Mask coefficients are generated in parallel to Protonet. Being an anchor-based detector, it has two branches in the prediction head, namely classification and bounding-box offset regression. To compute mask coefficients, another branch is added, in parallel to the other two. In the final step, the masks are assembled by combining the prototype branch and the mask coefficient branch. The operations are implemented as a matrix multiplication and sigmoid:

$$M = \sigma(PC^T)$$

where $P$ is a $h \times w \times k$ prototype masks and $C$ is a $n \times k$ matrix of mask coefficients corresponding to $n$ instances that passed through Non Maximum Suppression (NMS). In addition to the new architecture, they also propose an improved version of NMS, called FastNMS, where the decision of keeping or not an instance is done in parallel for all instances, therefore improving the speed performance even more. Although this version suffers from removing a little bit too many boxes, the accuracy drop is negligible in comparison to the huge increase in speed performance.

3.5. Particle Detection Using ConvNets

Object detection, instance segmentation and semantic segmentation techniques have been used in a variety of real-world applications, ranging from autonomous driving, aerial navigation to more industrial oriented applications, such as smart manufacturing or specialized tasks in factories. Therefore, these techniques have also been used for
estimating the PSD in different industrial applications and have added new challenges for these algorithms, pushing the level of innovation and demanding more research. It comes as no surprise that methods based on DL are proving to be more efficient than traditional methods even for this type of estimation, but they are relatively new and ongoing research is still needed.

In [44], authors use DL for estimating the distributions of grain size and porosity from micro-CT images. As training data, they generate synthetic 3D images of spheres, simulating how a micro-CT image would look like. They’ve chosen a 3D CNN [45], which was initially used for human action recognition from video images. The 3D CNN takes as input the 3D images and outputs directly two values, the grain size and the porosity label, hence in this case, a regression rather than segmentation is performed. Training was done on the synthetic data and then the model was tested on real-world data, showing promising results.

In the case of 2D images, we can also encounter the problem of partially sintered and agglomerated particles, since the 3D particles are projected on the image plane. [7] aims at addressing this problem by using Mask R-CNN architecture to first segment instances of particles and to compute the PSD. The same strategy is applied as in the latter article, training and validating on synthetic images and then testing on real-world scanning electron microscopy images (SEM) or transmission electron microscopy images (TEM). After inference on the test images, the PSD is calculated using the Feret diameter as the equivalent diameter of a particle. Although the method produces satisfying results in terms of accuracy, the time and memory requirements are not taken into consideration and may pose a problem if this solution would be integrated in a real-time application.

Another domain that picked the interest of deep learning and particle detection is Cryo-electron microscopy (cryo-EM). Here, detection is difficult mainly because of extremely low signal-to-noise ratio (SNR). The approach used in [46] is quite unique and uses two DNNs, one classification network and one segmentation network. First, the classification network is trained and then using its parameters as initial values for the segmentation network, the training process is accelerated. The segmentation network is responsible for getting probability (density) maps that are fed to the selection algorithm (Grid-based Local-maximum selection) and produces initial results. Ultimately, preliminary results are fed to the classification network and final results are obtained. Similar to [47], they use ‘Atrous convolution’ feature in the segmentation network. In terms of training data, because of the low SNR which causes very difficult manual annotation, they generate images from real-world datasets and also use simulated datasets.

Xiao Y. proposed in [48] a solution for particle picking based on the Fast R-CNN framework [35]. To develop a fast method, they have tried to solve one of Fast R-CNN’s bottlenecks, which was the region proposal. Instead of using selective search, they have proposed a sliding window approach. Their solution reduces the test time from 1.5 minutes obtained from [49] to 2 seconds. They have also tried to use Faster R-CNN, which replaced selective search with RPN, but argued that for cryo EM images, the RPN performs terrible and because of their special case where possible particles are of fixed size, the sliding window is much faster and better. Also, the papers discussed so far dealt with particle picking when there is only one particle of interest in the image, but in this case, their images contained ice particles, which makes classification harder,
specifically the rate of false positives increases. To deal with that, they have annotated ice particles as well and formulated the problem as a three-class classification, so that the neural network would learn the subtle difference between ice and protein particles. As a result, the rate of false positives decreased significantly.
4. DETAILED DESIGN AND IMPLEMENTATION

The focus of this thesis is building a system capable of estimating the PSD of mineral rocks from images taken in a flow-through system. From a commercial point of view, an emphasis is put on apatite particles, since they are the most valuable, but as a challenge and to ensure generality, we take into consideration phlogopite particles as well. For estimating the size distribution, we need to segment instances of particles in each image. Although this can be achieved with conventional image processing techniques, deep learning techniques prove to be much more robust, accurate and faster, representing the state-of-the-art in the instance segmentation task. Moreover, although semantic segmentation networks, like DeepLab [47], can also segment the image, the dataset on which we are working includes a lot of particles that overlap and would thus require post-processing steps that would slow the pipeline. Hence, we choose two instance segmentation networks, Mask R-CNN [38] and Yolact [43], both having their own specific advantages. Mask R-CNN represents the state-of-the-art architecture in instance segmentation, but Yolact is able to achieve real-time performance with sacrificing just a fraction of accuracy. An overview of the pipeline applied in this thesis is depicted in Figure 17. Regarding implementation, Matterport’s implementation [50] of Mask R-CNN has been used and for Yolact [43], authors have released their open-sourced repository.

4.1. Imaging Setup

This section will present the setup used for acquiring the data. Outotec’s PSI 500 was the device of choice, being capable of computing the size distribution using laser diffraction, hence having a reference measurement. A flow through cuvette was connected with a tube to PSI 500 outlet, such that the minerals are imaged in a flow-through system. There were two setups of lightning, front lightning which emphasized the color of particles and back lightning, which resulted in grayscale images. For our case, we have decided to use front lightning. The camera used for capturing images from the flow-through system had a magnification factor of 3.5 micrometers.
per pixel. The magnification factor was important for aligning the size distribution obtained from images with the real-world scale and also to compare it with the size distribution obtained by PSI 500. The dataset consisted of colour images, having a resolution of 2448x2048. Fig. 18 shows an example of captured images.

(a) Raw image data.  
(b) Annotated image.  
Figure 18. Images captured by PSI 500 and annotated with VIA tool.

4.2. Task Description

For this use case, there are 2 particles of interest, namely apatite and phlogopite. By looking at Fig. 18, one can already start discriminating between them. Apatite particles are more transparent and clear while the phlogopite particles are mostly opaque and have a brownish tint. The central task of this project is getting the size distribution of apatite particles across a sample of images. The stakeholders are mainly interested in apatite detection because phlogopite is much less valuable, therefore it is not so relevant for the grinding process. On the other hand, training an algorithm to detect phlogopite as well, may result in increased accuracy and may serve well for other use cases, where multiple different particles may need to be detected. Hence, the task is now formulated as a two-class instance segmentation problem.

The dataset consists of hard and tricky cases as well, where particles can be occluded, partially captured in the image and even particles that are mixed or hard to classify because of similarity. When particles are occluded or partially captured in the image, it represents a problem for the PSD because we do not know its actual size, hence we may alter the accuracy of the PSD. In our case, it was decided that particles laying on the border of the image can be discarded, provided that we have enough detected particles so that the result is statistically meaningful. Mixed particles are also interesting, because one can be interested in detecting the ratio of apatite/phlogopite from the particle or simply label it as mixed. There are also difficult particles to classify even for a human operator. Some can be semi-transparent and have a faint tint of brown, so it may be really difficult to assign one class. Another major challenge for this dataset is represented by making the distinction between a particle that is in-focus and out-of-focus. Because every measurement that it is performed needs to be converted to a real-world scale, for particles that are not in-focus we cannot approximate correctly their true size. The magnification factor is known only for the
region where particles are in-focus, everything else cannot be estimated by a simple linear adjustment. Therefore, only particles that are in-focus need to be detected, so we can get a reliable PSD. In Fig. 18, particles of interest are highlighted and annotated manually.

An important note is the lack of 100% correct ground-truth, since the provided dataset does not include annotations as well, hence particles need to be manually annotated, with limited knowledge in the domain of mineral processing. The only reference measurement available is the PSD provided by PSI 500, which unfortunately cannot discriminate between apatite and phlogopite and computes the PSD considering both particles and using the technique of laser diffraction.

4.3. Dataset Description

The datasets of images were provided by Outotec. There are 5 datasets, each having a different ratio of apatite and phlogopite particles, measured by another system. For each of the dataset, a reference measurement was provided by using the PSI 500 particle analyzer, namely a graph consisting of particle size (µm) and their frequency, which can be seen in Figure 24. Because the data had to be manually labelled, only one of the dataset was labelled and the others were used only for running the inference on the trained model and getting the particle distribution for comparison with the reference measurement. The only difference between the datasets was the distribution of apatite vs. phlogopite particles. As a result of the similar structure between datasets, it allows for easy transfer learning. For training a model, the dataset would need to be split in a training, validation and test set. A proportion of 60-20-20 was chosen, having 72 training images, 30 validation images and 33 test images.

4.3.1. Annotation Process

When dealing with DL models, one of the most common problems is data. For a model to be able to learn and to generalize well, a relatively big dataset is needed. There are a number of big datasets on the internet, like COCO [42], ImageNET [15], Cityscapes [51] and so on, but they cannot contain all the images from all domains, but they are rather intended as natural images, containing everyday scenes/objects. Hence, in the case of real-world data and specialized domains, researchers have to manually label their specific dataset, which in many cases is a laborious work. Also, depending on the task, from object detection to instance segmentation, the annotation can be more and more challenging. In the case of image classification, it can be fairly quick to label images, but going to instance segmentation, where one has to draw polygons carefully, it can take significant time, especially if we have lots of instances per image, and we need to label hundreds of images. For labelling the data, the VIA labelling tool [52] has been used, where one can draw polygons around the object of interest. Generating bounding boxes afterwards is really easy. The software exports the annotations in JSON format and can be directly used in Matterport’s implementation. When training the Yolact model, the implementation expects COCO-style annotations, therefore a script for converting the VIA annotations to COCO annotations was implemented.
When annotating the images, although the annotation task is fairly easy, few challenges were encountered. First of all, there are no clear guidelines to distinguish between the apatite or phlogopite particles. As a rule of thumb, the apatite particles are clear and transparent, while the phlogopite ones are less transparent, sometimes opaque and have a brownish color. Unfortunately, there are also particles that are mixed and contain both apatite and phlogopite, or the brownish color cannot be clearly seen. All of these are increasing the complexity of the annotation process.

Another challenge involved the annotation of particles that are not in-focus. The reason for not wanting particles out of focus is that after the segmentation, the particles need to be measured and then multiplied with the magnification factor of the camera. Particles that are not in focus do not have the same magnification, hence including out of focus particles will alter the accuracy of the PSD. Because of the small focus area, it was very hard in some cases to distinguish between what is in focus and what is out of focus. In some cases, large particles were partly in focus and partly out of focus because of their size. Another difficult case is when the particles are overlapping. Not only do they influence the color of one another, but sometimes it can be hard to correctly guess and label its true contour.

4.3.2. Dataset Analysis

Matterport’s implementation of Mask R-CNN offers scripts for analysing the dataset, which helps in carefully setting some of the network’s configuration parameters. The first analysis is carried on the mean value of all the pixels in the dataset, for each color channel. This value is useful for mean subtraction, as part of the network preprocessing steps. This also had to be done for Yolact as well. If we are dealing with images of multiple dimensions, we also need to analyse their dimension and choose the optimal size, as input images are resized to one size, so the model can be trained with multiple images per batch. Other statistics, such as number of particles per image or per dataset can be obtained, which can be useful for the evaluation of the network. A function for computing the bounding box distribution was also added, as this allows for better choosing the anchor scales. The anchors can be overlaid on the input images, so we can hint how they are covering the feature maps. Unfortunately, Numpy uses bytes for Boolean values, resulting in large masks for high resolution images, making the training really slow. For this, mini masks are used (resize masks to a smaller scale e.g. $56 \times 56$). They can also be inspected to see which mask size achieves the optimal trade-off between accuracy and low memory.

The size of particles was also analyzed by plotting the histogram of equivalent diameters. Figure 19 reveals that most of the particles have the equivalent diameter less than 290 pixels, which translates to roughly 1 mm in metric scale. The maximum size that can be detected by PSI 500 is 1 mm, but by looking at the histogram, the imaging system allows for identifying even larger particles.
4.3.3. Transfer Learning

As mentioned above, the size and quality of data play a major role in building a good DL model. State-of-the-art DNNs require thousands of labelled images, which in some cases, as in this work, is just impossible to get.

Transfer learning paradigm [53, 54] hopes to offer a solution to this problem by transferring the common features that are shared by multiple data points. This technique consists of training a model on a big dataset, like ImageNet or COCO, and use the weights as initial weights for training the model on a much smaller, specific dataset. The reasoning behind why this works is that many images share the same low-level spatial characteristics, and it is much easier for the model to learn these features from big data. Not only it solves the problem of having a small training dataset, but transfer learning is also a technique used for preventing overfitting. In this work, pre-trained weights of models trained on COCO and ImageNet have been used, but the difference in accuracy between them is negligible considering the use case of this thesis.

Depending on the implementation and models, in some cases a restrictive rule is to have the same image size as the images on which the pre-trained model was trained. In the case of Matterport’s implementation, that was not an issue, but in the case of Yolact, only 550x550 or 700x700 images can be used. Therefore, that means scaling down the images from 2448x2048 to 550x550, which greatly affects the quality of the images and annotations. Especially smaller particles became so small that it was impossible to visually localize them, while the blurriness was accentuated a lot. Moreover, rescaling the dataset makes the comparison between models a bit more difficult and inconsistent.

![Particle size histogram for the training dataset.](image)
4.3.4. Data Augmentation

Transfer learning solves the problem of having huge data for training, but it works until a certain moment. While it can transfer low-level spatial features, we still need to deal with the high-level ones. And even for this task, the available dataset was really small, namely only 73 images for training. Data augmentation is the technique used to artificially generate new, similar data from existing one. The aim of data augmentation is to increase the size of a dataset. There are two major categories of image data augmentation according to [55], specifically basic manipulations and deep learning approaches. In this work, the focus will be on the first one, consisting of basic image processing techniques that alter the images in a slight way, so that the dataset size is increased with plausible fabricated images. This process is applied only to the training dataset as opposed to data preparation, where the same operations need to be applied to the validation and testing dataset as well. The choice of image processing operations needs to be taken carefully, by analyzing the context of the data. For example, it does not make any sense to rotate a picture with a car upside-down as the model will most surely not see anything like that.

For the training dataset, in the case of Mask R-CNN, the considered augmentations were flipping of the image left/right, flipping up/down, rotating it, blurring, sharpening and edge enhancement. With each new augmentation operation, the dataset doubles its size. For the current use case, a lot of flipping and rotations can be done, since spatially, the particles can be anywhere in the image and placed in any direction. But when dealing with colors, it was essential to capture the brownish color of the phlogopite as a distinctive landmark, consequently no operations altering the color channels has been performed. If using augmentations that change the spatial location of the objects, annotations also have to be modified accordingly. Yolact data augmentation consists of photometric distortion, random sample crops, mirroring, flipping and rotations.

4.4. Training and Optimization

When training a DNN, the two most important metrics are training and validation loss. They represent all the losses summed up and offer a general overview of the network performance and how well it is learning. The losses have to be evaluated together, so that we can choose the best model. By analyzing the training loss, we can see if the model has started to learn or not, by how the loss converges or spikes. But a converging training loss does not indicate the performance of the model and from a certain point, the model actually overfits the training data, resulting in poor generalization. The validation loss can indicate the overfitting, as it stops improving after a certain amount of epochs and afterwards it keeps degrading. Validation loss is also used for early stopping [56], namely stop training after there is no more improvement in the validation loss for a specified interval.

In Fig. 20, the training and validation loss from the Mask R-CNN model are shown. The Matterport’s implementation saves a new model whenever a better validation loss has been recorded, so at the end of the training we are sure to get the best model. Unfortunately, Yolact’s implementation does not provide such a feature, but by analyzing the losses, we can select the best model.
In Mask R-CNN, we are training the model with the full-resolution images, so that we can accurately segment small particles as well. Because of this, we can work with a batch size of only one image. On the other hand, on Yolact we have to work with much lower resolution images ($550 \times 550$) and the architecture is more lightweight, meaning that we can have a batch size of even 8 images, which is highly recommended by the authors. A bigger batch size usually allows for a better convergence to the local minimum when applying gradient descent.

Other parameters, such as weight decay, learning rate, learning momentum, etc. were left the same as in the original implementation. Adjusting different parameters and obtaining different models can easily become hard to track and then especially difficult to compare them. Also, visualizing independently the losses (class loss, bounding-box loss, mask loss) can yield some insights into what may go wrong with the training. A great tool for solving all these issues is W&B (Weights and Biases) [57]. With few lines of code, it is able to log all the desired metrics and stores them in the cloud, offering a nice centralized visualization. This way, we can compare the losses between different models and choose which one was the best one and what parameters were used. It can also keep track of all the losses, so they can be inspected separately.

### 4.4.1. Blurry Class

When doing the evaluation of the first model, we observed an interesting fact: the rate of false positives was significantly high, and they usually occurred when the model was detecting out-of-focus particles. Strictly considering detection, this proves how remarkable the network is and how capable it is of detecting a good variety of particles. On the other hand, considering the use case of PSD, out-of-focus particles represent false positives, and they should be avoided. Our idea was to formulate again the problem, but as a three class instance segmentation problem, adding the blurry class, which would have blurry apatite and phlogopite particles under the same category, simply blurry particles. This way, when running the inference on the data, we get in the same time in-focus and out-of-focus particles. There are also cases in which a particle is detected both as an apatite/phlogopite and as a blurry particle, but considering the NMS, the detection with higher confidence is kept.
4.5. Size Distribution

After training the network, we obtain the best model, which will be used for inference on the unlabeled datasets. Once the particles are segmented, we can easily compute their area in terms of number of pixels. To be able to compare the results with the ones obtained from laser diffraction, we need to approximate the area by the equivalent diameter, which is done using:

\[ D = 2 \cdot \sqrt{\frac{A}{\pi}} \]

where \( D \) is the diameter and \( A \) is the area of the particle expressed in pixels. The final step consists in converting from pixels to real-world measurement, by applying the magnification factor of the camera. For the camera in our experiment, the magnification is 3.5 micro-meters per pixel.

There are other methods as well for measuring the size distribution, especially suited for when particles have shapes that are more elongated. In [58], the authors use the Feret’s diameter, which is the longest dimension of the particles, independent of its angular rotation.
5. TESTING AND VALIDATION

The quality of the estimated PSD is heavily dependent on how well the instance segmentation task is performed. The first requirement that needs to be satisfied is the accuracy of our models. Because instance segmentation is a more complex task, we need to combine multiple metrics to get a complete understanding of how these models perform and to evaluate them accordingly. Working with closed-source datasets from customers means that evaluating a model requires more in-depth analysis. The results obtained on the current datasets cannot be straightforward compared to other open-source datasets, simply because the data is significantly different. Most researchers develop their DNN architecture and benchmark it against well-known open-source datasets, such as COCO [42], PASCAL [59], CityScapes [51] etc. These datasets contain general objects which are clearly distinguishable and in-focus. On the contrary, our dataset exhibits some difficult challenges, considering that even for a human operator it may be difficult to discriminate between apatite or phlogopite or in-focus vs. out-of-focus. Therefore, the metrics are applied on data that has been manually annotated and considered as ground-truth by non-experts. While these metrics would validate the model on the instance segmentation task, the PSD computed from the instance segmentation can be assessed by comparing it to the PSD obtained by Outotec’s PSI 500 device. The second requirement that needs to be satisfied after ensuring accuracy is real-time performance. PSD is relevant only if we have a large enough number of samples, and it is updated in real-time, so that we can infer the result on the whole population. For our setup, this can be achieved if the processing time is at least 10 FPS.

The purpose of the evaluation phase is to assess the performance of DNNs in the context of instance segmentation applied on mineral images and to compare the PSD obtained by DL-based segmentation with the one resulted from PSI 500, which relies on laser diffraction. This chapter first introduces and defines the metrics that will be used in the evaluation phase, presenting the obtained results on our specific dataset. Then, the size distribution results are revealed and analysed in comparison with the reference measurement provided by Outotec. Ultimately, size distributions obtained by the two models are compared against one another.

5.1. Metrics

To measure the performance level of DNNs with the intent of comparing them, we use a range of metrics. Especially when dealing with complex tasks such as instance segmentation, simply looking at the classification accuracy of the model is not enough. Popular metrics include confusion matrix, precision-recall curve (PR curve) and mean average precision (mAP) score. The metrics are performed for both Mask R-CNN and Yolact. We also measure the inference time in frames per second and consider it a vital part in comparing the two architectures from the overall system’s point of view.
5.1.1. Confusion Matrix

As its name suggest, the confusion matrix is a matrix that expresses how many instances we managed to detect correct, incorrect or could not detect at all. In literature, the following terms are used:

- true positives (TP): the model correctly predicts an instance
- true negative (TN): the model correctly predicts a misdetection. This does not apply to object detection or instance segmentation, since it would represent all bounding boxes (masks) that are not detected in an image, which would be a huge number.
- false positives (FP): the model incorrectly predicts an instance
- false negatives (FN): the model is not capable to predict an instance

Computing these terms gives a better image of how the model is performing and what should be improved. But in object detection or instance segmentation, the classification of a detection is not binary, like detected or not detected, as the detection can partially overlap with the ground-truth. Because of this, another metric is needed, to establish what we consider a valid or not valid detection.

Intersection Over Union (IoU) is a measure based on the Jaccard Index that evaluates the overlap between two bounding boxes or instances, known as detection and ground-truth. The formula for IoU score is depicted in Fig. 21. The IoU score is a number between 0 and 1 and represents how much the two bounding boxes are overlapping. By setting a threshold (usually 0.5), a detection can then be classified as a TP or FP and subsequently for a ground-truth instance, we can tell if it is a FN or not.

\[
\text{IoU} = \frac{\text{area of overlap}}{\text{area of union}}
\]

Figure 21. Intersection over Union mathematical formulation.

In our experiments, we benchmarked the two models by using the test dataset. The validation dataset is used only in the training phase, to be sure that there is no bias towards the data on which the evaluation metrics are calculated. Confusion matrices for both Mask R-CNN and Yolact models have been generated and the results are depicted in Table 1. To have a fair comparison, we generated the confusion matrices by adopting the same thresholds, setting the confidence score threshold at 0.3 and the IoU threshold score at 0.5. Predictions that have the confidence score below the threshold
are discarded, as well as predictions that have lower IoU score than the threshold are considered as two separate instances.

<table>
<thead>
<tr>
<th></th>
<th>Yolact</th>
<th>Mask R-CNN</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Apatite</td>
<td>Phlogopite</td>
</tr>
<tr>
<td>True Positives</td>
<td>112</td>
<td>117</td>
</tr>
<tr>
<td>False Positives</td>
<td>155</td>
<td>42</td>
</tr>
<tr>
<td>False Negatives</td>
<td>20</td>
<td>59</td>
</tr>
</tbody>
</table>

Table 1. Performance metrics for test data.

The test dataset consists of 132 apatite particles and 176 phlogopite particles. By focusing on the Mask R-CNN results only, the number of TP is exceptionally high while the number of FN is low, meaning that the model can detect the majority of apatite particles. In the case of phlogopite particles, the number of undetected particles is higher, but still at a decent level. However, the number of FP is much higher than TP, both for apatite and phlogopite particles. After visually inspecting the inference on the test images and comparing it with the ground-truth, the following conclusions have been drawn. First, the network is able to detect exceptionally small particles, which were ignored in the annotation process. On the other hand, it has problems in distinguishing between in-focus and out-of-focus particles, detecting many particles that are blurry. If only the instance segmentation task is considered, then the network performs really well in finding all the particles, but in our specific use case, blurry particles are not relevant.

Relating the results from Mask R-CNN to Yolact, we can draw the conclusion that the first model is capable of detecting more particles. Although the number of FP is relatively low compared to that of Mask R-CNN, a potential explanation is the rescaled images to lower resolution for Yolact model, which makes the detection of really small particles almost impossible.

The confusion matrix for both models shows that the model is detecting too many particles and the visual inspection confirms the fact that the models are struggling in avoiding blurry particles. As a potential solution, we have decided to label particles, both apatite and phlogopite that are blurry, in the hope that the system will learn the difference between in-focus apatite and phlogopite and blurry particles. After blurry particles have been identified, they are no longer used in the size distribution estimation. The NMS is also modified to favour blurry particles, by suppressing other detections if the current one is a blurry one with the confidence score above 0.5. The results are documented as a confusion matrix in Table 2.

<table>
<thead>
<tr>
<th></th>
<th>Yolact</th>
<th>Mask R-CNN</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Apatite</td>
<td>Phlogopite</td>
</tr>
<tr>
<td>True Positives</td>
<td>110</td>
<td>118</td>
</tr>
<tr>
<td>False Positives</td>
<td>159</td>
<td>39</td>
</tr>
<tr>
<td>False Negatives</td>
<td>22</td>
<td>58</td>
</tr>
</tbody>
</table>

Table 2. Performance metrics for test data. The model was trained with the blurry class.
Despite lowering a bit the number of FP, there is an increase in FN. The same applies for both architectures. Although the confusion matrix is a robust performance measurement, one has to fix a confidence score threshold and an IoU score threshold, which offers just a snapshot. Optimal thresholds can be chosen by applying a metric that is based on varying these scores and assess how the model performs.

5.1.2. Precision-Recall Curve

Precision - Recall measure (PR) is a useful performance measure when the classes are imbalanced and when the main objective is to find a trade-off between parameters. Precision represents the rate of FP, while recall represents the rate of FN. They can be mathematically formulated as:

\[
P = \frac{TP}{TP + FP} \\
R = \frac{TP}{TP + FN}
\]

We can assess the performance of the model by visually inspecting the area under the PR curve. A high area indicates high precision and high recall, meaning low FP and FN rates. A system with high precision, but low recall returns few results, but most of them are correct. On the other hand, a system with high recall, but low precision returns many results, but most of them are incorrect. The ideal system has both high precision and recall, but in real-world scenarios, we have to aim for a trade-off between these two, depending on the use-case of the system.

Figure 22 shows a comparison of PR curves between two models, both trained on Mask R-CNN architecture, but one has been trained with an additional blurry class for blurry particles. On the same figure, the IoU score threshold is varied and different confidence scores are highlighted.
Following the same procedure, we compute the PR curves for the Yolact model, trained with and without the blurry particle class. Figure 23 shows the obtained results. For Yolact, it is obvious that the addition of the third class, blurry particle, results in an improved precision and recall. To achieve a trade-off between precision and recall, the best configuration would be an IoU score of 0.5 and a confidence score of 0.5. Depending on the specificity or sensitivity requirements, different configurations can be selected. Unexpectedly, for certain configurations, the Yolact model proves to work even better than the Mask R-CNN, precisely in terms of precision, since Yolact is much less likely to predict false positives.

Figure 22. Precision-Recall curve for test set using Mask R-CNN.

Figure 23. Precision - Recall curve for the test set using Yolact model.
Mean average Precision

Most of the time though, expressing the overall performance in just a number is desired due to its simplicity and ease of understanding. The Average Precision (AP) represents one of the most used metric for assessing a DL model in the task of instance segmentation. The AP is defined as the area under the PR curve.

\[ AP = \int_0^1 p(r) \, dr \]

As a pre-processing step, usually the PR curve has a zig-zag pattern and needs to be smoothed out. This is done by replacing each precision value with the maximum precision value to the right of that recall score. Then, an interpolation is performed to get the final result. In COCO, a 101-point interpolation is performed on the PR curve. Traditionally, AP is performed across a range of IoU thresholds (0.5-0.95). The mean Average Precision (mAP) is the average of AP's calculated at all IoU thresholds between 0.5 and 0.95, for all classes.

5.1.3. Inference Time

To have a complete analysis of the two DNN architectures applied for this use case, a comparison in terms of processing speed must be performed. Both architectures had a ResNet 101 backbone. Regarding image size, Mask R-CNN was trained with 2048x2048 images, while for Yolact, images had to be resized to 550x550, which resulted in slight accuracy loss, but gained significant processing time. Even if the input image size differs, according to [43], Yolact is supposed to be faster than Mask R-CNN. In our experiments, inferencing on a sample of images resulted in 23 FPS processing speed for Yolact and only 5 FPS for Mask R-CNN. The requirement for online grinding control and monitoring is a processing time of at least 10 FPS.

5.2. Size Distribution

All the evaluated metrics above rely on one essential concept: having a reliable, fixed ground-truth. As it was specified in Section 4.3.1, the annotation process is not straightforward and can suffer from errors due to the highly specialized domain. Because of this, it is mandatory to evaluate our models with respect to a reference measurement, which does not rely on manual annotation. Outotec provided reference measurements by operating the PSI 500. Figure 24 depicts the size distributions for different datasets, by applying the laser diffraction method. Due to the inherent inability of laser diffraction to distinguish between particles of different classes, the DL-based pipeline must also combine the size distributions obtained from the apatite and phlogopite particles in a mixed distribution, for achieving a correct comparison.
Figure 24. Reference measurements provided by Outotec’s PSI 500, using laser diffraction technology.

Figure 25 shows the PSD of all samples, when Mask R-CNN model has been used, in two formats: histogram and kernel density estimation (KDE). The process of obtaining a histogram for a specific dataset is presented in Section 4.5. The same process is applied for all samples and each histogram is normalized individually. However, plotting all histograms on the same canvas leads to cluttered areas, from which it can be hard to analyse the result. Additionally, due to the frequent spikes in the histograms, a smooth curve that approximates the underlying distributions would be desirable. This can be achieved by calculating the KDE for each sample, which replaces the discrete histogram with a continuous, smooth curve that characterizes the observed PSD. It can also introduce potential distortions, if the underlying data is not smooth enough or bounded, but choosing a good set of smoothing parameters can alleviate this problem. The computation of KDE was performed with the help of Seaborn [60], a Python data visualization library, that uses only Gaussian kernels. The final curves are plotted on a logarithmic scale, so that the axes match with the reference measurement.

When comparing the reference PSD with the ones obtained by our method, we have to take into consideration certain aspects. First, the laser diffraction method is capable of measuring only particles of up to 1 mm, which translates to approximately 286 pixels. By referring to Section 4.3.2 and Figure 19, the imaging system is capable of detecting larger particles, hence the size distribution might vary from the reference measurement. Second, after a visual inspection of the datasets, it was observed that a significant number of medium-large particles, usually phlogopite, are out-of-focus, due to the narrow depth-of-field. Hence, there is a strong possibility that such particles are measured by the laser diffraction, but for the current imaging system it is impossible to correctly process them.

Analysing Figure 25 and Figure 24, it seems like sample4 and sample5 still have the highest magnitude and sample1, sample2 and sample3 follow the trend of two spikes, although the magnitude is reversed. Comparing the positions of distributions, DL-based system detects particles that are smaller than particles detected by the laser diffraction. Surprisingly, the PSD obtained when applying Yolact indicates different distributions. Corroborating the shape and position of the PSD curves with confusion matrix and with the limitation of Yolact processing only 550x550 images, it becomes
clear that Yolact model is having difficulties in detecting smaller particles. The shapes of PSD curves are tighter than that of the ones obtained with Mask R-CNN, while the peaks are slightly shifted to the left.

(a) Histogram of particle size  
(b) Kernel density estimation for the PSD

Figure 25. PSD estimated by Mask R-CNN model trained with blurry labelled particles.

(a) Histogram of particle size  
(b) Kernel density estimation for the PSD

Figure 26. PSD estimated by Yolact model trained with blurry labelled particles.

To better understand the impact of each model on the PSD, Figure 27 shows the ground-truth PSD compared to the PSD obtained by the two models. The graph supports the hypothesis that Mask R-CNN is more biased towards detecting small, blurry particles than Yolact is. Furthermore, the inference PSD results validate that DL models can successfully segment instances of particles and produce PSDs similar to the ground-truth. We have also analysed the PSD of apatite and phlogopite individually by plotting the graph in Figure 28, and the same conclusion was reached, that small apatite particles are detected by the Mask R-CNN. Regarding phlogopite, the PSDs look more similar and closer to the ground-truth, since they are easier to segment.
Figure 27. PSD of ground-truth instances and the PSDs resulted from inference with Mask R-CNN and Yolact.

Figure 28. Individual PSD for apatite and phlogopite.

A shortcoming of the laser diffraction method is that it cannot distinguish between different particles, hence the resulting size distribution is a characteristic of the mixture of particles. Whereas if using the imaging system, particles can be identified separately and size distribution can be likewise computed individually or for the whole mixture. In Figure 29, we have plotted the histogram of apatite and phlogopite particles and computed the KDE for one of the samples.
Figure 29. Histogram and KDE for apatite and phlogopite particles of sample 1.
6. DISCUSSION

This work investigates the application of DL-based instance segmentation in the context of estimating PSD. To achieve this goal, two DNNs, specialised for the task of instance segmentation, have been trained with images of particles in fluid suspension. The first architecture to be tested was Mask R-CNN, a robust and powerful architecture, especially suited for transfer learning. Although the model yielded good results on our dataset, the major drawback was the slow processing time and considering the purpose of the overall system, which is real-time particle monitoring, we had to shift the focus towards an architecture more specialized in real-time processing. Yolact is designed to be a simple, lightweight, real-time DNN for instance segmentation, while maintaining a high level of accuracy. An extensive comparison was performed between the two models to understand the key differences and to obtain an optimal trade-off between accuracy and speed, so that the requirements for an online particle monitoring system would be fulfilled. Moreover, the PSDs obtained from the DL-based pipeline were compared to a reference measurement provided by the PSI 500 particle analyzer from Outotec, based on the laser diffraction principle. Despite implementing a successful proof-of-concept, there are still a number of challenges that need to be addressed in order to have a robust and reliable mechanism for PSD estimation.

Inspired by the emerging concept of transfer learning, utilizing DNN for specialized data and use cases seems like a promising idea. The known phrase that ‘a DNN is as good as the data you feed it’ has solid foundations and applies for each DL architecture. Nevertheless, in specialized domains, good data is tremendously difficult to obtain and even much harder it is to label it correctly and efficiently. Thanks to transfer learning, we can utilize the general knowledge of a network to solve our particular task. This results in significantly less labelled data and reduced training time. Transfer learning represents the core concept of this work. The obtained results support the idea of transferring knowledge, since the models are pre-trained on huge datasets containing general objects and then the last layers of the models are fine-tuned for our specific dataset.

Unfortunately, there is still a significant part of manual annotation involved. One potential solution inspired from I. Yalniz et al. [61], would be to employ an iterative pipeline that would expand the training set. In the beginning, few labelled instances would be needed so that the system can be trained. Then, unlabelled data would be fed to the network and only predictions that have a high confidence score would be retained. Next, we would use the new labelled data by our model to expand the training set and we would re-train the model, with the purpose of achieving better accuracy. There are few investigations of how such frameworks would work, but it seems like a promising new concept.

The dataset proved to be complex and required careful analysis. Blurriness has also been a major challenge when carrying out the experiments. First, it is hard to distinguish between an in-focus and out-of-focus particle, especially because of the slurry composition that surrounds the particles. Apatite particles are significantly more difficult to recognize as blurry due to their transparent nature. Second, some particles, especially the large ones, are big enough to be partly in-focus and blurry at the same time. This is caused by the narrow depth-of-field, resulting in possible inaccurate
measurements if such particles are considered. It is also not clear yet if the DNNs can learn to distinguish between blurry and in-focus particles. The results obtained from training with a blurry class support the idea that the DNN is capable of identifying some of the blurry particles, but it is not yet robust enough. It may be that the network has to be modified in order to also calculate a blurriness measure and to be part of the learning and inference process, rather than a post-processing step, so that the impact on the inference time is not significant. Moreover, comparing the PSDs from the PSI 500 with the ones obtained from the imaging system can be almost impossible, as the laser diffraction can take into account also the blurry particles. Another observation resulted from the annotation process was that some particles can be also mixed, consisting of both apatite and phlogopite. Also, some cases were extremely hard to manually annotate, namely attributing a class, due to increased similarity in particles. In turn, this can affect the evaluation, as annotation is biased by the human annotator, especially if he or she is not an expert in the mining field.

To overcome the detection of small particles in the case of Yolact, a possible solution is to partition the original sized image, for example in four equal patches, then run inference on each of them individually and in the last step, assemble the final picture. The complex part lies in stitching back the four patches and solving the case of particles that are split between the patches. Naturally, this operation would add an overhead for the processing time, but may increase the number of small detected particles.

An experiment utilizing controlled particle size distributions obtained by sieving enriched material was planned to strengthen the evaluation part of this thesis. This would have allowed further analysing the performance of the approach. The experiment would have been conducted first with pure apatite, allowing analysing multiple distributions and directly comparing to the laser diffraction result. Then, material from the same distribution would have been mixed with phlogopite to evaluate the capability in a real scenario. Unfortunately, because of bad circumstances leading to delays in the project, such an experiment was not feasible anymore.

There are still areas that need further research in order to apply DNNs to real-world, hard engineering problems, but as the work performed up to this point shows, the trend looks promising.
The principal objective of this thesis was the study of DL-based instance segmentation for obtaining the PSD. Obtaining the PSD in real-time serves for control and monitor processes in the grinding circuit. The dataset comprised of images containing apatite and phlogopite particles in fluid suspension, captured by a color camera. The particles had to be identified in images and for that, two DNN models, specialized for the task of instance segmentation, were trained and used for inference on the unlabeled data. Once the particles have been identified, the PSD can be easily computed.

In contrast to other works [7], not only testing, but also validation and training is performed on real-world data by leveraging the concept of transfer learning. No synthetic data is generated and the data is manually annotated, taking in consideration difficult cases where particles are mixed, partially blurred or overlapping with other particles. The first trained architecture was Mask R-CNN, which provided exceptional results, unfortunately being too slow for real-time processing. The focus was shifted towards Yolact, a DNN designed with real-time performance in mind. In agreement with the author’s claims, the results of applying Yolact confirmed both the real-time processing and high accuracy, by achieving more than 20 FPS and comparable accuracy to Mask R-CNN. Although a reference PSD was provided, the measurement mechanism relies on laser diffraction, which operates entirely different and comparing with the PSD obtained with the imaging system can be really difficult.

However, a key advantage of the image-based system, corroborated with the high precision and real-time processing of the DNN, consists of computing a separate PSD for each particle of interest. In our case, the apatite PSD is of utmost importance and the proposed system is capable of completing this task. Moreover, it is capable of distinguishing between blurry and in-focus particles and even detect overlapped particles. The potential of using DNN in PSD estimation still needs to be explored and refined further. Furthermore, limiting the amount of manual annotation is one of the biggest challenge and this will be the focus of our future research.
8. REFERENCES


