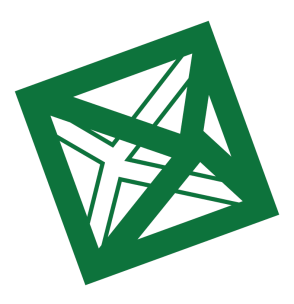
Università degli Studi di Milano-Bicocca

School of Science
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Master Degree in Particle Physics

Overcoming medical data scarcity: transfer learning from synthetic particle jets images to lung CT using deep neural networks

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Abstract

Early diagnosis is the most effective weapon in the fight against cancer, with computed tomography and medical imaging playing a crucial role in this process. Recently, artificial intelligence has shown great potential to identify and classify tumor lesions in their early stages. However, the application of these techniques faces a fundamental limitation: the scarcity of annotated data available for training.

As is well known in machine learning, the quality of results depends strictly on the quality and completeness of the training data ("garbage in, garbage out"). To overcome this limitation, this thesis explores the use of transfer learning, a technique that allows the transfer of knowledge from a data-rich source domain to a data-limited target domain.

Specifically, the research focuses on the use of simulated images of high-energy particle jets, constructed specifically for this study, characterized by two distinct classes of physical phenomena: a noisy background with physical characteristics similar to the main jets and morphological properties similar to tumor structures.

The proposed approach involves a pretraining phase on a large high-energy physics dataset, transferring the learned weights to the medical domain, and then fine-tuning them on limited clinical datasets. This method aims to leverage the feature extraction capabilities developed in the particle physics domain, adapting them to the medical context where annotated data are scarce, but diagnostic accuracy is crucial.

Chapter 1

Computed Tomography

Computed Tomography (CT) is a medical imaging technique that uses X-rays to generate detailed cross-sectional images of the human body. Since its introduction in the 1970s, CT has revolutionized diagnostic radiology by enabling the visualization of internal structures with high spatial resolution and three-dimensional reconstruction capabilities. Unlike conventional radiography, which produces a single projection image, CT acquires multiple X-ray measurements from different angles around the patient and uses computer algorithms to reconstruct the corresponding anatomical slices.

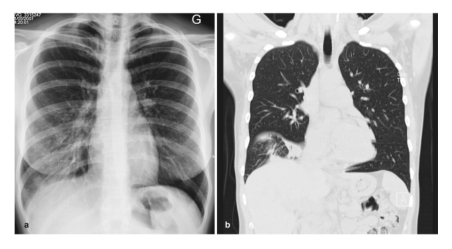


Figure 1.1: On the left (figure a.) we can see a chest radioraphy, on the right (figure b.) we can see a CT scan of the same area. The CT scan provides a much more detailed view of the internal structures, allowing for better diagnosis and treatment planning.

CT plays a crucial role in clinical practice due to its ability to provide fast, accurate, and non-invasive diagnostic information for a wide range of applications. Modern CT scanners are equipped with advanced technologies that significantly improve image quality, reduce scan time, and minimize radiation dose.

This chapter provides an overview of the main aspects of CT imaging that are relevant for medical physics applications. The first part briefly describes the experimental setup of a CT scanner, the second part focuses on the mathematical principles behind image reconstruction, highlighting the algorithms used to convert projection data into tomographic slices. Finally, the chapter introduces the concept of image post-processing, with particular attention to windowing and its impact on image interpretation.

1.1 Tomography setup

The basic idea behind tomography is to obtain a three-dimensional image of an object from a series of two-dimensional images acquired from different angles. For this purpose, a CT scanner consists, like any radiation measurement system, of a radiation source (an X-ray tube) and an array of particle detectors. To acquire projections from multiple angles, typically ranging from 0 to 180 degrees, the X-ray tube and the detectors are mounted on a rotating gantry, which allows the system to rotate around the patient or, more generally, around the object being examined.

As shown in Figure 1.2, the detectors are not arranged on a plane perpendicular to the X-ray source, but rather along a circular arc. This configuration ensures that all pixels in the projection correspond to the same physical size, thus avoiding geometric distortions in the reconstructed image. Depending on the scanner design, the acquisition can be performed in a step-and-shoot mode or using continuous rotation, as in helical (spiral) CT. The X-ray beam is collimated into a fan or cone shape to cover the entire detector array. Each detector element typically consists of a scintillator coupled to a photodiode, which converts the X-ray photons into an electrical signal proportional to the received dose. Synchronizing the gantry rotation with data acquisition allows the collection of a sufficient number of projections to ensure accurate image reconstruction.

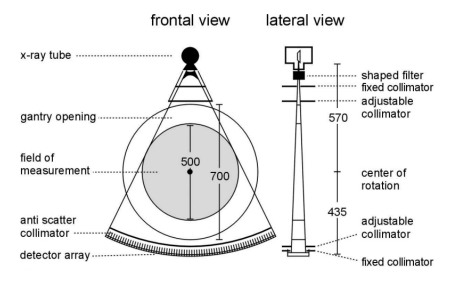


Figure 1.2: In the frontal view we can see, in order, the X-ray tube (radiation source), the rotating gantry (support for the X-ray tube and detectors), the field of measurement (where the patient or the object is allocated), the anti scatter collimators (to reduce the amount of scattered radiation reaching the detectors), and the detectors (which measure the intensity of the X-rays after they pass through the object). In the lateral view we can see the presence of collimators, used to adjust the narrowness of the beam depending on the type of scan and the area of interest.

1.2 Image Reconstruction

In computed tomography (CT), image reconstruction refers to the process of transforming raw projection data, acquired from different angles, into a cross-sectional image of the scanned object. The core idea is to recover a 2D function f(x,y) — representing the

internal structure — from its projections. Each projection contains information about how much an X-ray beam is attenuated as it travels through the object along a certain direction.

1. The Radon Transform

Mathematically, each X-ray beam travels along a straight line. For a beam oriented at angle θ , the line is defined by:

$$x\cos\theta + y\sin\theta = s$$

where s is the perpendicular distance from the origin to the line.

The projection $p(s, \theta)$ is obtained by integrating the object's density f(x, y) along that line:

$$p(s,\theta) = \int_{-\infty}^{+\infty} f(x,y) \, \delta(x \cos \theta + y \sin \theta - s) \, dx \, dy$$

This is called the **Radon transform** of the function f(x, y).

2. The Fourier Slice Theorem

To reconstruct the image, a key result is the **Fourier Slice Theorem**. It says that the 1D Fourier transform of a projection at angle θ gives a slice of the 2D Fourier transform of f(x,y) taken in the same direction.

We compute the 1D Fourier transform of the projection:

$$P(\theta, u) = \int_{-\infty}^{+\infty} p(s, \theta) e^{-2\pi i u s} ds$$

Plugging in the expression for $p(s, \theta)$, we get:

$$P(\theta, u) = \iint f(x, y) e^{-2\pi i u(x \cos \theta + y \sin \theta)} dx dy$$

This expression matches the 2D Fourier transform of f(x,y), evaluated at:

$$u_x = u\cos\theta, \quad u_y = u\sin\theta$$

so we can write:

$$P(\theta, u) = F(u \cos \theta, u \sin \theta)$$

In other words, each projection gives us a radial slice of the 2D Fourier transform of the image.

3. Inverse Transform and Filtering

If we collect projections over many angles, we can build the 2D Fourier space of the image. Then, to reconstruct f(x, y), we apply the inverse 2D Fourier transform:

$$f(x,y) = \iint F(u_x, u_y) e^{2\pi i (u_x x + u_y y)} du_x du_y$$

Switching to polar coordinates:

$$u_x = u\cos\theta, \quad u_y = u\sin\theta$$

the Jacobian of the transformation gives:

$$du_x du_y = |u| du d\theta$$

Substituting into the inverse Fourier formula:

$$f(x,y) = \int_0^{\pi} \int_{-\infty}^{+\infty} |u| P(\theta, u) e^{2\pi i u(x\cos\theta + y\sin\theta)} du d\theta$$

Here, the factor |u| acts as a high-pass filter, enhancing high-frequency components and correcting for the blurring introduced during projection. The equation above gives the foundation for the **Filtered Back Projection** algorithm, one of the most common reconstruction methods in CT. It works as follows:

- 1. Acquire projections $p(s, \theta)$ at many angles $\theta \in [0, \pi]$.
- 2. Compute their 1D Fourier transforms $P(\theta, u)$.
- 3. Multiply by |u| (or an equivalent ramp-like filter) to enhance sharpness.
- 4. Invert the Fourier transform to get filtered projections.
- 5. Back-project the filtered projections over the image domain and sum them.

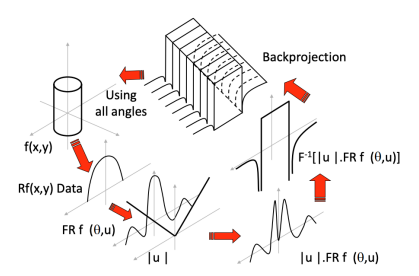


Figure 1.3: Radon transform and reconstruction via filtered back projection. Each projection contributes a slice in frequency space; the image is recovered by inverse Fourier transform.

The filtered back projection is fast and effective, which is why it is still widely used in clinical CT. However, modern scanners often integrate more advanced techniques such as iterative reconstruction or machine learning, which can offer better noise suppression and lower radiation dose. Nevertheless, understanding the Radon transform and the Fourier Slice Theorem provides deep insight into how tomographic images are formed.

1.3 Windowing

The pixel values are quantified in **Hounsfield units (HU)**, a standardized scale that reflects the radiodensity of the tissue. This scale is defined relative to the attenuation coefficients of water and air:

$$HU_{tissue} = 1000 \times \frac{\mu_{tissue} - \mu_{water}}{\mu_{water} - \mu_{air}}$$
 (1.1)

where μ represents the linear attenuation coefficient. Key reference points: water = 0 HU for definition, air = -1000 HU.

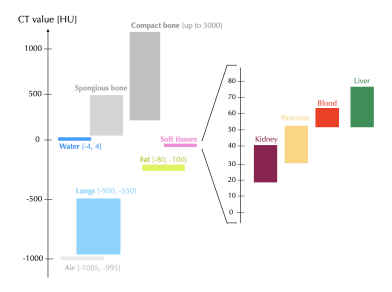


Figure 1.4: CT Hounsfield Unit (HU) ranges for key tissues. Highlighted regions show typical HU values for diagnostic reference, from dense compact bone ($\sim 3000~{\rm HU}$) to air (1000 HU), with soft tissues (20 - 80 HU) and fluids (blood $\sim 45~{\rm HU}$, water 0 HU) in intermediate ranges.

Windowing is a technique used in computed tomography (CT) to enhance the visibility of specific tissues or structures within the body. It involves adjusting the range of Hounsfield units (HU) displayed in the CT image, allowing radiologists to focus on particular types of tissues, such as bones, soft tissues, or air-filled spaces. The window width and center determine the contrast and brightness of the image, respectively.

- Window width (WW): This parameter controls the range of Hounsfield units displayed in the image. A narrow window width enhances the contrast between tissues with similar densities, while a wider window width allows a broader range of densities to be visualized.
- Window center (WC): This parameter sets the midpoint of the Hounsfield unit range displayed in the image. Adjusting the window level changes the brightness of the image, allowing radiologists to focus on specific tissue types.

The choice of window width and level depends on the specific clinical question and the type of tissue being examined.

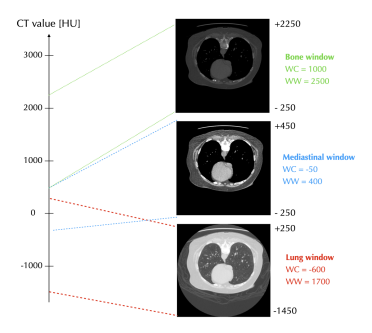


Figure 1.5: Standard CT windowing settings for different anatomical structures. Full CT value range ([-1000, 3000] HU). Clinical presets with window center (WC) and width (WW) values for bone (WC=1000, WW=2500), mediastinum (WC=-50, WW=400), and lung (WC=-600, WW=1700) visualization.

1.4 CT Planes

Images can be visualized in three standard orthogonal planes: axial, coronal, and sagittal. Each plane offers a unique perspective on anatomical structures, aiding in comprehensive clinical assessment.

- Axial plane (transverse): A horizontal plane that divides the body into superior (upper) and inferior (lower) parts. It is the primary acquisition plane in most CT scans.
- Coronal plane: A vertical plane that divides the body into anterior (front) and posterior (back) parts. Often used for viewing the frontal anatomy.
- Sagittal plane: A vertical plane that divides the body into left and right portions. It is particularly useful for evaluating asymmetries between the two sides.

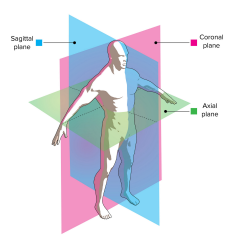


Figure 1.6: CT planes: axial, coronal, and sagittal. The axial plane is horizontal, the coronal plane is vertical and divides the body into front and back, and the sagittal plane is vertical and divides the body into left and right.

Chapter 2

Machine Learning in a nutshell

Machine learning is a subfield of artificial intelligence that focuses on the development of algorithms and statistical models that enable computers to perform tasks without explicit instructions. Instead, these systems learn from data, identifying patterns and making decisions based on the information they have been trained on. Machine learning consists of designing efficient and accurate prediction algorithms. More generally, learning techniques are data-drivem methods combining fundamental concepts in computer science with ideas from statistics, probability and optimization.

Types of tasks

The following are some standard types of tasks in machine learning:

- Classification: Assigning labels to data points based on learned patterns (e.g., e-mail spam detection).
- Regression: Predicting continuous values based on input features (e.g., predicting house prices). In regression, the penalty for an incorrect prediction depends on the magnitude of the difference between the true and predicted values, in contrast with classification problem, where there is typically no notion of closeness between various categories.
- Object detection: Identifying and localizing objects within images or videos (e.g., detecting tumours in lung CT scans).
- Clustering: Grouping similar data points together without predefined labels.
- **Anomaly detection:** Identifying unusual patterns that do not conform to expected behavior.
- Ranking: Ordering items based on their relevance or importance.

Algorithms that solve a learning task based on semantically annotated historical data are said to operate in a **supervised learning** mode. In contrast, algorithms that use data without any semantic annotation are said to operate in an **unsupervised learning** mode. In the latter case, the algorithm is expected to discover patterns in the data without any prior knowledge of the labels or categories. In this thesis I'll mainly focus on supervised learning.

Label set: We use Y to denote the set of all possible labels for a data point of a given learning problem. Note that the labels can be of two different types: categorical

labels, which are discrete and finite and define classification problems, and continuous labels, which can take any value in a continuous range and define regression problems.

2.1 Neural networks and deep learning

Neural networks are a class of machine learning algorithms inspired by the structure and function of the human brain. They consist of interconnected nodes (neurons) that process information in layers. Deep learning refers to the use of neural networks with many layers (deep neural networks) to model complex patterns in large datasets. This approach has led to significant advancements in areas such as image recognition, natural language processing, and game playing.

2.2 YOLO: You Only Look Once

Object detection is a task that involves identifying and classifying objects present in images or videos. Initially, object detection was approached as a pipeline consisting of three main steps: proposal generation, feature extraction and region classification. However, this approach was computationally expensive and often led to suboptimal results. The emergence of deep learning brought a significant change in object detection, with deep convolutional neural networks (CNNs) playing a crucial role in this transformation. CNNs are designed to automatically learn hierarchical features from raw pixel data, eliminating the need for manual feature engineering. This shift allowed for more efficient and accurate object detection systems.

Currently, deep learning-based object detection frameworks can be classified in two families:

- Two-stage detectors: These methods first generate region proposals and then classify them. Examples include R-CNNs (Region-based Convolutional Neural Networks), that first generate region proposals using a selective search algorithm and then extracts features from these regions using a CNN; the extracted features are then fed into an SVM for object classification.
- One-stage detectors: These methods perform detection in a single pass, directly predicting bounding boxes and class probabilities. Examples include YOLO (You Only Look Once), which exists in eleven versions. The YOLO models are popular for their accuracy and compact size. It is a state-of-the-art model that could be trained on any hardware. YOLOv8, in particular, was developed by Ultralytics and introduced on January 2023. It is used to detect objects in images, classify images and distinguish onjects from each other.

2.2.1 Architecture of YOLOv8

The YOLOv8 architecture is composed of two major parts, namely the **backbone** and the **head**, both of which use a fully convolutional neural network.

• The **backbone** is responsible for extracting features from the input image. It consists of a modified version of the CSPDarknet53 architecture, which has 53 convolutional layers and employs a technique called cross-stage partial connections to

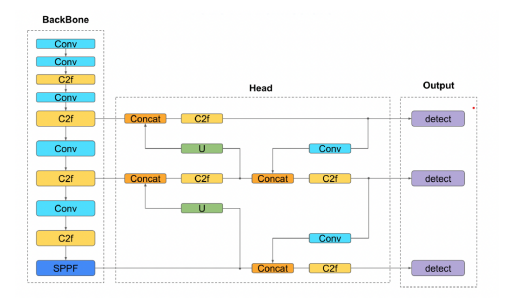


Figure 2.1: Architecture of YOLOv8. The backbone extracts features from the input image, while the head predicts bounding boxes and class probabilities.

enhance the transmission of information across the various levels of the network. The convolutional layers are organized in a sequential manner to extract relevant features from the input image.

• The **head** is responsible for predicting bounding boxes and class probabilities. It consists of a series of convolutional layers that take the features extracted by the backbone and apply additional operations to predict the bounding boxes and class probabilities for each object in the image. The head uses a technique called anchor boxes to handle objects of different sizes and aspect ratios.

The YOLOv8 framework can be used to perform computer vision tasks such as detection, segmentation, classification and pose estimation and comes with pre-trained models for each task. For detection, the models are pre trained on the COCO dataset, while for classification on ImageNet dataset. There are different versions of YOLOv8, each designed for different tasks and with different architectures. The most common versions are YOLOv8n, YOLOv8n, YOLOv8m, YOLOv8l and YOLOv8x, where the letter indicates the size of the model (n for nano, s for small, m for medium, l for large and x for extra large). The larger the model, the more parameters it has and the more computational resources it requires to train and run.

2.3 Object Detection Metrics

A metric is a function that quantifies and evaluates the performance of a model. In the context of object detection, metrics are used to assess how well a model can detect and classify objects in images.

Intersection over Union (IoU)

When we talk about object detection, we often refer to the concept of **bounding boxes**. A bounding box is a rectangular box that is used to define the location of an object in an

image, typically represented by its top-left corner coordinates (x, y), width, and height, or top-left and bottom-right coordinates. The bounding box is used to localize the object within the image and is often used in conjunction with a classification label to identify the object. Intersection over Union (IoU) is a metric used to evaluate the accuracy of the bounding box detection. It measures the overlap between the predicted and the ground truth box. The IoU is calculated as follows:

$$IoU = \frac{Area_{Intersection}}{Area_{Union}} = \frac{A_{pred} \cap A_{gt}}{A_{pred} \cup A_{gt}}$$
 (2.1)

The IoU value ranges from 0 to 1, where 0 indicates no overlap and 1 indicates perfect overlap. A common threshold for considering a detection as a true positive is an IoU of 0.5 or higher.

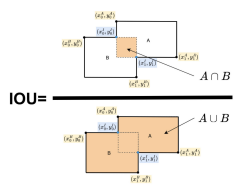


Figure 2.2

Algorithm 1 Intersection over Union (IoU) between two bounding boxes

- 1: **Input:** Boxes $A = (x_A, y_A, w_A, h_A)$ and $B = (x_B, y_B, w_B, h_B)$
- 2: Compute bottom-right corners:

$$x2_A = x_A + w_A, \quad y2_A = y_A + h_A$$

 $x2_B = x_B + w_B, \quad y2_B = y_B + h_B$

3: Compute intersection rectangle:

$$x_{\text{int}}^1 = \max(x_A, x_B), \quad y_{\text{int}}^1 = \max(y_A, y_B) x_{\text{int}}^2 = \min(x_{A}, x_{B}), \quad y_{\text{int}}^2 = \min(y_{A}, y_{B})$$

4: Compute intersection area:

$$w_{\text{int}} = \max(0, x_{\text{int}}^2 - x_{\text{int}}^1)$$

$$h_{\text{int}} = \max(0, y_{\text{int}}^2 - y_{\text{int}}^1)$$

$$A_{\text{int}} = w_{\text{int}} \times h_{\text{int}}$$

5: Compute areas of each box:

$$A_A = w_A \times h_A, \quad A_B = w_B \times h_B$$

6: Compute union area:

$$A_{\rm union} = A_A + A_B - A_{\rm int}$$

$$A_{
m union} = A_A + A_B - A_{
m int}$$
7: Return: $IoU = \frac{A_{
m int}}{A_{
m union}}$

Precision and Recall

Precision and recall are fundamental metrics used to evaluate the performance of object detection models. These metrics provide valuable insights into the model's ability to identify objects of interest within images. Let's define some fundamental concepts before diving into precision and recall:

- True Positive (TP): These are istances where the model currently identifies and localize objectys, and the intersection over union (IoU) score between the predicted and the ground truth bounding box is equal or greater then a specified treshold.
- False Positive (FP): These are cases where the model incorrectly identifies an object that does not exist in the ground truth or where the predicted box has an IoU score below that define treshold.
- False Negative (FN): These rapresent instances where the model fails to detect an object that is present in the ground truth. This means that the model has not detected an object that is present in the image, resulting in a missed detection.
- True Negative (TN): Not applicable in object detection. It represents correctly rejecting the absence of objects, but in object detection, the goal is to detect objects rather than the absence of them.

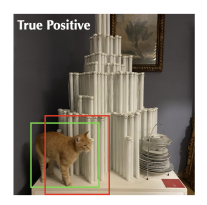


Figure 2.3: The object is there, and the model detects it, with an IoU above treshold.

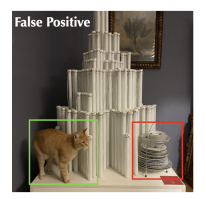


Figure 2.4: The object is there, but the predicted box has an IoU below the treshold.



Figure 2.5: The object is there, and the model doesn't detect it. The ground truth object has no prediction.

Figure 2.6: Ground truth box is the green rectangle, while the predicted box is the red rectangle.

Let's not define the core metrics:

• **Precision**: Is a critical metric in model evaluation as it serves to quantify the accuracy of the positive predictions made by the model. It specifically assesses how well the model distinguishes true objects from false positives. In essence, precision provides insight into the model's ability to make positive predictions that are indeed accurate. A high precision score indicates that the model is skilled at avoiding false positives and provides reliable positive predictions.

$$Precision = \frac{TP}{TP + FP} \tag{2.2}$$

• Recall: also known as sensitivity or true positive rate, is another essential metric used in evaluating model performance, especially in object detection tasks. Recall measures the model's capability to capture all relevant objects in the image. In essence, recall assesses the model's completeness in identifying objects of interest. A high recall score indicates that the model effectively identifies most of the relevant objects in the data.

 $Recall = \frac{TP}{TP + FN} \tag{2.3}$

We can define the **Average Precision (AP)** as the area under the precision-recall curve, which is obtained by plotting precision against recall for different confidence thresholds ¹. The AP is a single value that summarizes the model's performance across different levels of precision and recall. It is calculated as follows:

$$AP = \int_0^1 P_{\alpha}(box_{gt}, box_{pred}) dR_{\alpha}$$
 (2.4)

where P_{α} is the precision at a given IoU threshold α and R_{α} is the recall at the same threshold. Precision signifies the accuracy of the model's positive predictions, while recall quantifies the model's ability to successfully identify all relevant objects. AP achieves a harmonious balance between false positive and false negatives, providing a comprehensive evaluation of the model's performance.

Mean Average Precision (mAP)

Mean Average Precision (mAP) extends the concept of Average Precision (AP) by providing a global summary of a model's performance across multiple object classes and, in some cases, multiple localization thresholds. While AP evaluates the precision-recall trade-off for a single class — by varying the confidence threshold and computing the area under the resulting precision-recall curve — mAP averages these AP values to reflect the model's overall detection capabilities.

$$mAP = \frac{1}{N} \sum_{i=1}^{N} AP_i \tag{2.5}$$

where N is the number of classes and AP_i is the Average Precision for class i.

In practice, several versions of mAP are commonly reported. mAP@0.5 refers to the mean AP calculated at a fixed Intersection over Union (IoU) threshold of 0.5. mAP@[0.5:0.95], computes the average AP across ten IoU thresholds ranging from 0.5 to 0.95 (in increments of 0.05), providing a more rigorous and comprehensive evaluation that accounts for both classification and localization accuracy.

$$mAP@50 = \frac{1}{N} \sum_{k=1}^{N} AP_{K}@50$$
 (2.6)

¹The confidence threshold is a tunable parameter used to filter out low-confidence predictions in object detection tasks. Each predicted bounding box is assigned a confidence score, typically representing the estimated probability that the detection both contains an object and correctly identifies its class. Varying this threshold affects the trade-off between precision and recall: lower thresholds increase recall but may introduce more false positives, whereas higher thresholds improve precision at the cost of reduced recall.

$$mAP@[50:95] = \frac{1}{N(95-50)} \sum_{\alpha=50}^{\alpha=95} \sum_{k=1}^{N} AP_{K}@\alpha$$
 (2.7)

where N is the number of classes and α is the IoU threshold.

2.4 Transfer learning

Transfer learning is a machine learning technique that leverages knowledge gained from one task to improve performance on a different but related task. It is particularly useful when the target task has limited labeled data, allowing models to benefit from pre-trained representations learned from larger datasets. Transfer learning can be applied in various domains, including computer vision, natural language processing, and speech recognition.

2.5 Curriculum learning

Chapter 3

Hadronic Collisions and Jet Reconstruction

3.1 High energy collisions in hadronic accelerators

In a hadronic collider, such as the LHC or the Tevatron, beams of hadrons—primarily protons and antiprotons—are collided at extremely high energies. Unlike electrons, protons are not elementary particles; they possess a complex internal structure composed of quarks and gluons. This internal structure is described by the parton distribution functions (PDFs), denoted in Fig. 3.1 as $f_i(x_j, \mu_F)$, where i indicates the parton type (quark or gluon), and j refers to the beam. The PDF represents the probability of finding a parton of type i carrying a fraction x_i of the proton's momentum at the time of collision.

Therefore, when two protons P_1 and P_2 collide, the interaction does not occur between the protons themselves, but rather between their constituents—the partons—with momenta x_jP_j . The partons that do not participate in the hard scattering process are referred to as underlying events, as illustrated in the figure.

When high-energy quarks are produced in the final state, they emit additional partons through a cascading process known as parton showering. This process continues until the energy scale drops sufficiently, at which point the partons recombine into observable hadrons in a process called hadronization.

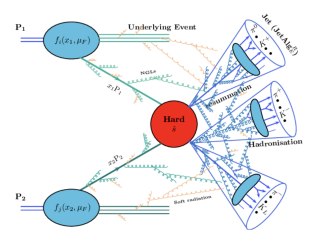


Figure 3.1: Schema di un collisione adronica. I protoni si scontrano e producono una cascata di particelle, che si trasformano in jet di particelle.

Jets

In order to reconstruct the primary event that occurred before the parton shower and hadronization, the strategy is to measure the energy of the final-state particles using calorimeters, and then group them based on their transverse momentum and direction. This procedure leads to the formation of what are known as jets. Jets are essentially the result of a clustering algorithm that groups final-state particles with similar transverse momentum and direction, allowing us to trace back to the primary parton-level event that initiated the particle cascade.

3.2 Useful variables to describe an hadronic process

In collider physics, the kinematic properties of particles are typically described using variables that are well suited to the cylindrical geometry of the detectors and to the characteristics of the collisions. Among these, the most commonly used are the transverse momentum p_T , the azimuthal angle ϕ , and the pseudorapidity η .

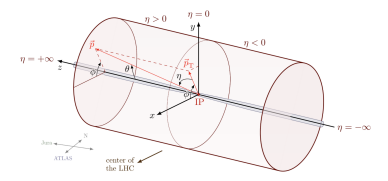


Figure 3.2: Kinematic variables used in collider physics. The transverse momentum p_T is the component of momentum perpendicular to the beam axis, ϕ is the azimuthal angle, and η is the pseudorapidity.

p_T - transverse momentum

The transverse momentum p_T is defined as the component of a particle's momentum that is perpendicular to the beam axis (commonly referred to as the z-axis in collider experiments). Mathematically, it is given by:

$$p_T = \sqrt{p_x^2 + p_y^2} (3.1)$$

where p_x and p_y are the components of the momentum in the plane transverse to the beam (also called the transverse plane).

This quantity is especially important in hadron colliders, where the incoming protons travel along the z-axis, but the exact longitudinal momentum of the interacting partons is unknown on an event-by-event basis. However, since the protons travel toward each other along the beam axis, the net transverse momentum of the system before the collision is essentially zero. As a result, conservation of momentum implies that the total transverse momentum of the final-state particles should also be zero, up to detector resolution and effects from invisible particles.

For this reason, p_T is a Lorentz-invariant quantity under boosts along the beam axis, and provides a frame-independent way to study the dynamics of the collision. It is also widely used in searches for new physics, where an imbalance in the total transverse momentum (often called missing transverse energy, or MET) can signal the presence of non-interacting or undetected particles.

ϕ - azimuthal angle

The azimuthal angle ϕ describes the direction of a particle in the transverse plane, i.e., the plane perpendicular to the beam axis (the x-y plane if the beam is aligned along the z-axis). It is defined as the angle between the particle's transverse momentum vector and a fixed reference axis (typically x-axis) measured counterclockwise:

$$\phi = \tan^{-1} \left(\frac{p_y}{p_x} \right) \tag{3.2}$$

The azimuthal angle takes values in the range $[-\pi, \pi]$ or $[0, 2\pi]$, depending on convention. This variable is invariant under longitudinal boosts.

η - pseudorapidity

The pseudorapidity η is a measure of the angle of a particle relative to the beam axis, defined as:

$$\eta = -\ln\left(\tan\left(\frac{\theta}{2}\right)\right) \tag{3.3}$$

where θ is the polar angle of the particle with respect to the beam axis.

3.3 Algorithms for jet reconstruction

Jet reconstruction algorithms are essential for identifying and grouping particles produced in high-energy collisions into jets, which represent the hadronic remnants of the original partons. These algorithms can be broadly categorized into two main types: **cone algorithms** and **sequential clustering algorithms**.

3.3.1 Cone Algorithms

Cone algorithms assume that particles in jets will show up in conical regions and thus they cluster based on $\eta - \phi$ space, resulting in jets with rigid circular boundaries. Most of the cone algorithms are iterative cones (IC). In such algorithms, a seed particle i sets some initial direction, and one sums the momenta of all particles j within a circle of radius R around i in azimuthal angle ϕ and pseudorapidity η , i.e. taking all j such that

$$\Delta R_{ij}^2 = (y_i - y_j)^2 + (\phi_i - \phi_j)^2 < R^2$$
(3.4)

where y_i and y_j are the pseudorapidities of particles i and j, respectively, and ϕ_i and ϕ_j are their azimuthal angles. The total momentum of the jet is then calculated as the sum of the momenta of all particles within this cone. The direction of the jet is then recalculated as the weighted average of the momenta of all particles within the cone, and the process is repeated until no more particles can be added to the jet.

3.3.2 Sequential Clustering Algorithms

Gli algoritmi a ricombinazione sequenziale utilizzano informazioni geometriche ed energetiche per aggregare le particelle che potenzialmente arrivano dallo stesso partone, ricostruendo così la storia del getto.

k_T Algorithm

The k_T algorithm is a sequential clustering algorithm that uses the concept of distance between particles to group them into jets. The distance between two particles i and j is defined as:

$$d_{ij} = \min(p_{T,i}^2, p_{T,j}^2) \frac{\Delta R_{ij}^2}{R^2}$$
(3.5)

while the distance between a particle i and the beam axis is defined as:

$$d_{iB} = p_{Ti}^2 (3.6)$$

For each particle, the algorithm calculated the minumum distance betwen d_{ij} and d_{iB} .

$$d_{min} = \min(d_{ij}, d_{iB}) \tag{3.7}$$

If d_{min} is smaller than a predefined threshold, the particle is added to a jet; otherwise, it is considered a standalone particle. The algorithm proceeds iteratively, recalculating the distances after each clustering step until all particles are grouped into jets or no more particles can be clustered.

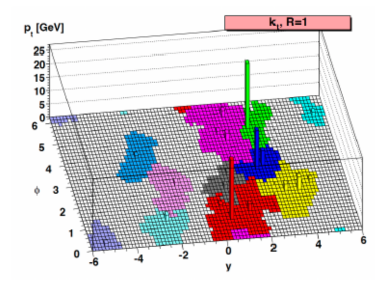


Figure 3.3: Schematic representation of the k_T algorithm. The algorithm iteratively clusters particles based on their transverse momentum and distance in $\eta - \phi$ space, forming jets.

Anti- k_T Algorithm

The anti- k_T algorithm is a variant of the k_T algorithm that uses a different distance measure to cluster particles. It is designed to produce jets with more uniform shapes and

is particularly effective in high-energy collisions where jets can be closely spaced. The distance between two particles i and j is defined as:

$$d_{ij} = \min(\frac{1}{p_{T,i}^2}, \frac{1}{p_{T,j}^2}) \frac{\Delta R_{ij}^2}{R^2}$$
(3.8)

while the distance between a particle i and the beam axis is defined as:

$$d_{iB} = \frac{1}{p_{T,i}^{2\beta}} \tag{3.9}$$

The algorithm proceeds similarly to the k_T algorithm, calculating the minimum distance between particles and the beam axis, and clustering them into jets based on this distance. The key difference is that the anti- k_T algorithm tends to produce jets with more uniform shapes, making it more suitable for high-energy collisions where jets can be closely spaced.

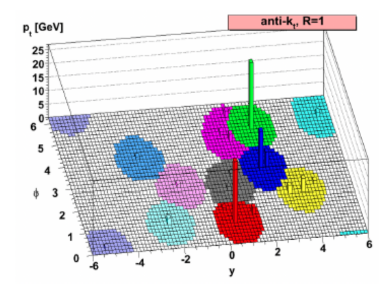


Figure 3.4: Schematic representation of the anti- k_T algorithm. The algorithm iteratively clusters particles based on their transverse momentum and distance in $\eta - \phi$ space, forming jets with more uniform shapes.

Cambridge/Aachen Algorithm

The Cambridge/Aachen algorithm is another sequential $\operatorname{anti} k_T$ clustering algorithm that uses a different distance measure to cluster particles. It is designed to produce jets with more uniform shapes and is particularly effective in high-energy collisions where jets can be closely spaced. The distance between two particles i and j is defined as:

$$d_{ij} = \Delta R_{ij}^2 \tag{3.10}$$

while the distance between a particle i and the beam axis is defined as:

$$d_{iB} = 1 \tag{3.11}$$

Physically, the differences between the three algorithms are contained in the momentum weighting. For the k_T algorithm, the weighting is done to preferentially merge

costituents with low transverse momentum with respect their nearest neighbours. For the anti k_T , the weighting is done so as to preferentially merge constituents with high transverse momentum with respect to their nearest neighbours. The C-A algorithm relies only on distance weighting with no k_T weighting at all.

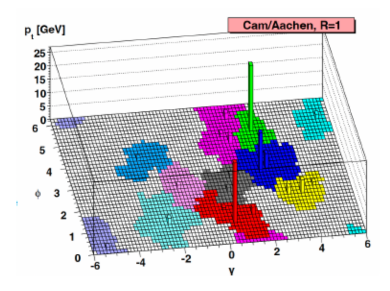


Figure 3.5

Chapter 4
Synthetic Particle Dataset

Chapter 5

Medical Dataset

5.1 NLST - National Lung Screening Trial Dataset

The aggressive and heterogeneous nature of lung cancer has thwarted efforts to reduce mortality from this cancer through the use of screening. The advent of low-dose helical computed tomography (CT) altered the landscape of lung-cancer screening, with studies indicating that low-dose CT detects many tumors at early stages. The National Lung Screening Trial (NLST) was conducted to determine whether screening with low-dose CT could reduce mortality from lung cancer. The NLST was a large, multicenter, randomized controlled trial that enrolled over 53,000 participants at high risk for lung cancer. Participants were randomly assigned to receive either low-dose CT scans or standard chest X-rays. The primary outcome was lung cancer mortality, with secondary outcomes including the detection of early-stage lung cancer and the impact of screening on overall mortality.

Radiologists at the screening centers reviewed the images obtained at each of the three annual screening exams to check for signs of lung cancer.

The image review was made without reference to any historical images. The radiologist recorded information about all visible abnormalities and assigned a preliminary screening result. A positive screening result (suspicious for lung cancer) was assigned if any non-calcified nodules or masses $\geq 4mm$ in diameter were noted or if any other abnormalities were judged suspicious for lung cancer by the radiologist.

5.1.1 Dataset Pruning

The NLST dataset contains chest CT scans where each scan comprises multiple (~ 143) axial slices. Notably, not all slices include tumor lesions. The dataset was then pruned by filtering slices based on a provided CSV file that identifies slices containing clinically relevant findings. After pruning, approximately 9,000 slices remain with malignant lesions with $\geq 4mm$ in diameter. However, benign tumor slices lack bounding box annotations, which presents a limitation for detection tasks.

The annotations are in the form of a CSV file that contains the following columns:

- **PID**: Unique identifier for each patient.
- Slice Number: The specific slice within the CT scan, which is also the name of the file.
- CT filter: Indicates the type of filter applied to the CT scan.

• x, y, width, height: Coordinates of the bounding box around the lesion, x and y represent the top-left corner of the bounding box, while width and height represent its dimensions.

5.1.2 Extraction of the pixel array

The CT images are in the dicom format, which is a standard format for medical imaging data. Each CT scan consists of multiple slices, and each slice is represented as a 2D image. The pixel values in these images are typically stored in Hounsfield units (HU), which represent the radiodensity of the tissue. A dicom file contains:

- Pixel Array: The pixel array is a 2D array of pixel values that represent the intensity of the image. Each pixel value corresponds to a specific location in the image and is typically stored as a 16-bit integer.
- Metadata: The metadata contains information about the image, such as the patient's name, the acquisition date, the image orientation, pixel spacing, and slice thickness. This information is essential for interpreting the image correctly.

5.1.3 Image Metadata

The metadata of a CT image contains important information about the image acquisition parameters and the physical characteristics of the image. The following are some key metadata fields commonly found in CT images:

- Pixel Spacing (mm): The physical distance between adjacent pixels in the x and y directions, typically measured in millimeters.
- Slice Thickness (mm): The thickness of each slice in the z direction, also measured in millimeters.
- Field of View (FOV) (cm): The physical dimension of the area captured in the image, typically measured in centimeters.
- Image Orientation: The orientation of the image, which can be specified using a combination of row and column vectors.
- Image Position: The position of the image in 3D space, typically specified using x, y, and z coordinates.

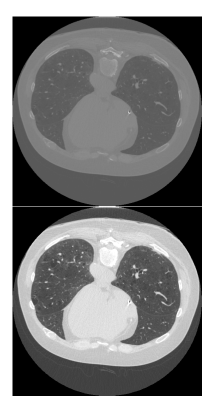


Figure 5.1: Didascalia principale delle due immagini affiancate

- 5.2 DLCS Duke Lung Cancer Screening Dataset
- 5.2.1 Extraction of the 2D slices
- 5.2.2 Dataset Pruning
- 5.3 Pre processing the images
- 5.3.1 Resampling
- 5.3.2 Windowing
- **RBG** Windowing
- 5.3.3 Padding
- 5.3.4 Masking