Chapter 2

Data Mining in Material Science

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Abstract
The invention of novel materials is the most propelling factor promoting the growth of contemporary civilization and technological innovation; nevertheless, previous materials research relied mostly on random techniques, which is arduous and labor-intensive. With the introduction of big data, which brings a deep upheaval in human society and considerably advances science, artificial intelligence, machine learning, and deep learning methods have recently made remarkable progress in materials science research. However, there are few comprehensive generalizations and descriptions of its applications in materials research. In this chapter, a brief summary of the evolution of materials science research is presented, followed by an emphasis on the key principles and basic processes of AI technique.

Keywords
Artificial intelligence, Machine learning, Deep learning, Materials

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1. Introduction

Artificial intelligence (AI) is a branch of computer science that aims to understand and build intelligent systems. Researchers have seen remarkable development in AI in recent decades, with some systems already achieving human-level and potentially super-human competence for a variety of activities such as voice recognition, picture analysis, machine interpretation, and games. There is a widespread conviction that AI will fundamentally alter many aspects of our culture and economy. Self-driving vehicles, for example, which incorporate actual visual identification and control, are on the verge of becoming a reality. This incredible development is causing a major transition in AI exploration from theoretical pursuit to an abundant larger sector with significant industry and governmental funding.

Considering the remarkable breakthroughs in AI, the research fraternity as a whole has taken notice and is investigating the application of AI for discoveries [1-11]. The materials science sector, in particular, has begun to use AI tools to speed material discovery. The current tendency is to apply machine learning (ML) approaches because many recent AI successes, particularly those involving superhuman powers are anchored in ML and deep learning (DL) approaches. Even though the conditions that facilitate DL are indisputable, particularly for visualization, language recognition, linguistic conversion, and independent driving, the constraints are soundly identified. In general, DL methodologies greatly rely on the accessibility of massive quantities of references or labeled data that is frequently

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unavailable. The present state of deep learning was likened to System 1. It comprises human's usual data processing because it is quick, easy, and a form of automatic pattern identification. Perception, encompassing vision, and hearing, is a component of System 1. The perceptual capacities are extremely advanced and a significant portion of the human intellectual cortex is dedicated to observation as depicted in figure 1.

Humans as well have an additional logical System 2, which is sluggish and requires significant thought and reasoning to solve difficult issues beyond reflexive reactions unlike System 1. Such complicated jobs are unsuitable for pure ML and DL. Nonetheless, AI spans a wide range of approaches, including hunting, cognitive, preparation, and data illustration. These strategies played a variety of parts throughout the AI growth stages. They are positioned to become more significant and play a growing role in supplementing pure ML techniques as well as further extending and propelling AI research to address increasingly complicated issues. These challenges include those involving scientific discovery, as well as jobs that need critical thought and reasoning and are carried out by people utilizing System 2.

![Figure 1 Structure of AI](image-url)

*Figure 1 Structure of AI [4]*
Before data-driven methodologies, material discovery evolved through three paradigms [12]. The first framework is based on experiments using trial approaches, the next is on the conversion of investigational data into a theoretical model, and the last is based on the computational simulation of theoretic representations. Individually model emerged in response to the outcomes of the preceding models. Recently, the fast expansion of existing data created through the preceding three models, along with the advent of the extremely effective algorithm, has permitted the use of AI, kicking off the quarter and the most current model of material discoveries. The main benefit of data-assisted and driven discovery is significantly reducing the necessary computing power, hence removing the limitations of the compositional transmission area. In recent years, artificial intelligence [10, 13-18] and virtual screening [19-27] approaches have been effectively applied to subfields of material and chemical science to identify novel organic compounds and constituents with extraordinary functions. Every area, due to its structural features, necessitates specialized approaches for simulated screening. Furthermore, to employ AI approaches in various sub-fields, a substantial quantity of great reliability and eminent data [28] must first be produced.

Two-dimensional materials (2D) are developing a sub-field of materials research. 2D materials, with their outstanding and controllable characteristics, show great potential for semiconductor, energy, and health applications [29-32]. Few unique 2D materials were fruitfully produced [33] later the finding of graphene [34], which has a meek 2D assembly of carbon atoms, however enticing and complicated physics. Despite a small number of experimentally produced compounds, novel 2D material depositories established on realistic considerable imitations have lately emerged [19, 35, 36]. The silicon repositories were created using two consecutive ways to build novel 2D materials: layer exfoliation through 3D bulk assemblies and combined atom interchange in 2D configurations. The initial method relies on the screening of exfoliated covered resources through 3D material databases [37-39]. This technique produces novel 2D material with assemblies that may utilize 2D patterns. The subsequent method encompasses exchanging single or additional atoms of an identified 2D pattern with molecules of extra chemical constituent. The following approach produces novel 2D constituents with identical crystal patterns as the original but with dissimilar chemical components. Regardless of the dual methodologies, the resource-intensive computation constrained the chemical search area for new 2D materials to the point where the quantity of 2D materials that were considered with density functional theory (DFT) calculations, the workhorse computational technique [40], has been reduced to a few thousand in recent years. Even though the DFT-calculated 2D materials records contain a few thousand materials, they are excellent equitable information repositories for data-driven algorithms. Amongst the primary aims, machine
learning is a promising way of exploring extraordinarily broad search spaces for possible chemical combinations in the order of particles, 2D and 3D materials. The procedure creates contestant resources systematically throughout an unparalleled chemical universe of molecules, recognizes probable stable resources, and forecasts important attributes of constant materials alternatives. They utilized this technique to create a computer-generated 2D Materials databank that comprises 316,505 probable constant 2D materials with projected attributes as a proof of concept [41].

2. Machine learning and materials science

“Machine learning” implementation approaches ML approaches which contain “supervised, unsupervised and reinforcement learning” and are an essential aspect of artificial intelligence. Each instance in supervised learning is made up of input and output, often referred to as a supervised signal. The labeling of data, comprising data grouping, information attributes, and feature idea placement, is one of the major aspects of supervised learning [42]. After training the algorithms with labeled information, factors of procedures are altered and predicted on the assessment outcomes of expected ones and predicted data and the process is repeated till the procedures congregate to optimal result, at which point the intelligent decision-making capability with the specific model is obtained. Unsupervised learning, unlike supervised learning, uses unlabelled data to search and derive probable links between samples. Clustering and dimensionality reduction are two common unsupervised learning methods. Clustering requires analysis of dispersal in feature areas of data measurements to categorize analogous information into single clusters and distinct information with dissimilar types. This is necessary because the categorization of data is unclear in advance. Furthermore, because higher dimensional information cliques are fairly uncommon in ML, difficulties such as distance computations and insufficient sample data are common, which is referred to as dimension catastrophe. The theoretical basis of reducing the dimension in ML is plotting the information sockets from novel higher dimensional to lower dimensional area and dimension reduction procedures primarily include factor analysis (FA), singular value decomposition (SVD), principal component analysis (PCA), and independent component analysis (ICA) [43-47].

Unlike “supervised and unsupervised learning, reinforcement learning” obtains knowledge statistics and updates the typical factors continuously based on input from the situation, it does not necessitate information to be supplied prior. In general, when the machine performs appropriately, a positive incentive is obtained; otherwise, a negative incentive is gathered. In this situation, the ML algorithms create several dynamic planning concepts, and eventually picked the action mode which maximizes the incentive. This demonstrates that the reinforcement learning technique requires less information and is easier to develop,
making it suitable for dealing with increasingly complex choice issues. Furthermore, as shown in figure 2 deep reinforcement learning that associates reinforcement through deep learning, is emerging as the fastest research hub in AI, particularly in automated driving, robots, language dispensation, and other domains [48-52]. Figure 3 depicts a framework of materials discovery which is used basically and designed based on ML approaches, with three primary processes listed: sample fabrication, algorithm model development, model verification, and materials prediction [53].

*Figure 2 Interlinking of AI, ML, and DL [53]*
Figure 3  Structural design of materials discovery and ML [53]
3. ML algorithms in materials science

The critical stage in the development of a machine learning system is the selection of an efficient machine learning system since it has a significant impact on generalization ability and prediction accuracy [55]. Every single algorithm has its unique set of applications; hence no approach is ideal for all problems. As seen in Figure 4, in materials science, the most often employed machine learning algorithms fall into 4 groupings: “probability estimation, regression, clustering, and classification” [54]. Probability estimate procedures are mostly utilized for novel material detection whereas “regression, clustering, and classification” techniques are primarily employed for predicting the material characteristics at all levels. Furthermore, ML approaches are frequently integrated with intellectual optimization techniques [56], which are primarily employed for optimizing model characteristics. Additionally, these techniques may be used to tackle additional complex tasks [57], like spatial configuration and material property optimization [58, 59] as portrayed in Figures 5 and 6.

Figure 4 ML algorithm in materials science [54]
Figure 5 Application of ML in materials science [54]
4. **Steps in machine learning for materials science**

Mitchell defines ML as a "mainframe program is supposed to acquire from experience E regard to a set of tasks T and performance measure P" [60] that is utilized as foundation background L in the following way as depicted in Figure 7.

4.1 **Experience**

Utmost ML algorithms may be classed as either unsupervised or supervised learning depending on the sort of samples that are provided throughout the training phase. Unsupervised learning uses a collection of instances containing just features to establish a few significant connections between the cases. A dataset of instances containing characteristics and associated labels that are “right” values linked with characteristics is used in supervised learning. Another type is reinforcement learning (RL), in which the representative knows through engaging with surroundings to acquire reward feedback. Conversely, RL confronts several obstacles, the most significant of which is the difficulty in creating interactive settings with rapid response. As a result, it is not yet commonly used in material discovery, and this evaluation confines itself to be guided and unsupervised.
### 4.2 Task

ML could be used for a variety of purposes. The task generally entails processing samples provided to an algorithm. The illustrations include features/descriptors that are typically organized as vectors to define the physicochemical characteristics, structural features, composition attributes, or formulation process conditions of the material.

### 4.3 Classification

The algorithm is concerned with determining which group or category a certain problem fits. It is accomplished through the training function \( f \) that transfers the characteristics vector to one of the unique modules. In its place of settling on a single group, \( f \) can instead provide a probability distribution across every group, with every element in the output vector representing the likelihood that the sample corresponds to the assured group. Material categorization challenges have been successfully solved using ML algorithms. A grouping model can forecast the effective formation of synthesized materials [61] and also the regions which exhibit any flow defect whenever given for specific synthesis parameters [62].

### 4.4 Regression

Regression is another typical job in which the algorithm attempts to study the function \( f = R_n/R \) that generates an uninterrupted \( y \) value or a series of data points represented as a vector \( y \). Regression models are commonly used to estimate material attributes like specific heat of objects, wave equation temperature, and optical gap of materials [63-65].

### 4.5 Clustering

It is a type of unsupervised learning technique, which comes in handy with a huge volume of unlabelled data [66-68]. It groups the set into clusters wherein elements inside the same group are more "similar" to one another than to another. What is meant by "similar" varies depending on context and circumstances. Even if the dataset is unlabelled, grouping objects into clusters allows for valuable insight into the data. Clustering algorithms have been used in material discovery in materials textual data mining and micro-structural image processing.

### 4.6 Dimension reduction and conception

While teaching a network, it may depend on several variables [69]. In such cases, it is advantageous to decrease the measurement of characteristics by projecting into smaller dimensions while retaining quite as abundant data as feasible [70]. It will assist in improving computational efficiency, and model performance, reduce overfitting, and aid
in the discovery of insights for specific applications. Dimension reduction was utilized to enhance prediction outcomes in material discovery, such as summarizing entire long unique features into lesser information to obtain better performance [71, 72]. Dimension reduction was employed to discover the essential science of physical prototypes by removing less important aspects [73]. Furthermore, mapping higher dimensional information to 2D or 3D charts for conception is a significant dimension reduction function since it allows meaningful insights to be drawn from an understandable graphic. It has been used to depict the high-dimensional material design space in material discovery[74].

4.7 Efficient searching

Conducting replications and lab-scale studies to collect more data is arduous and expensive. Effectual searching strategies can assist in identifying the best relevant additional data points to annotate, reducing total data-collecting efforts. Effectual probing was applied to the design of Perovskite, multilayer films, carbide, and nitride, among other materials [74, 75]. Overall, determining which objective to tackle is the initial stage in applying new ML in material discovery. Though, the five work types are not necessarily independent of one another.

4.8 Performance measure

The performance metric is employed to evaluate the performance of an algorithm on a certain job. The accuracy and error rate can be used to assess the algorithm's success in classification [76]. The log probability may be determined that produce probability dissemination. Mean square error or various variants of error are commonly used in the regression. A task's dataset is often divided into three parts: training data, a validation set of data, and data for testing. The algorithms will be taught during training and then fine-tuned based on their efficiency. The test efficiency of the optimized learning method is generally tested depending on the capability to execute test data, which functions as a measure of how effectively the model can generalize. The clustering technique in unsupervised learning may be quantified using exterior and interior indexes. Exterior indices require previously defined clustering configuration, i.e. the actual labeling for data in the dataset. The outcome of internal indices is assessed using quantities and attributes intrinsic to the database [77, 78].

4.9 Model particulars

The procedure of ML to employ is perspective and critical to select the proper model for a suitable task to obtain optimal effectiveness without inadequate or underfitting. Numerous ML models are often employed in the domain of AI-aided material discovery
and are described in many papers. These approaches are introduced in supervised, unsupervised, and weakly supervised learning paradigms. Figure 1 depicts the entire architecture.

4.10 Supervised model

The supervised learning algorithm is initially qualified using labeled data containing N training instances. The ith training example, for instance, consists of a combination of dual vectors for structures and tickets. The purpose of the model is to learn the given function. The model features are critical to be determined by the problem context. Figure 8 depicts a typical supervised learning approach in material discovery. It can create datasets through lab trials, simulations, or pre-existing datasets. There are two types of characteristics that are often employed in material discovery internal, and external information about the material classification. The intrinsic content is determined by the qualities of materials and chemicals utilized, but extrinsic data is determined by the surroundings wherein the materials reside. The two sorts of characteristics can be utilized as raw features directly. However, before feeding the raw characteristics into the prediction model, certain additional modifications are usually made to them. Normalization is the first pre-processing transformation that may be used to convert the quantities of attributes to a comparable basis. Another alternative pre-processing alteration is dimension reduction, which is employed if there are too many distinct features, particularly when compared to the number of labeled training samples. In this instance, it may be preferable to lower the dimension of the attributes to avoid overfitting. [14, 79]. In addition to the above-mentioned pre-processing methods, domain precise transition will be performed using the same preceding model. The preceding framework executes a few raw characteristics and transfers the output to the last forecasting framework. The translation of diverse chemical structures, for example, into fixed-length signature vectors can aid in improving forecasting ability in molecular property prediction tasks. Following these steps, the various treated and unrefined characteristics may be reduced to generate the complete characteristic given vector. Choosing the characteristics to employ is the utmost important stage in guaranteeing a successful model presentation [80]. Regardless of the highest excellent model, it will perform poorly if the features supplied to it are unsuitable. Following feature processing, several supervised models may be employed to forecast the finalized tasks. The most often employed supervised learning algorithms in material discovery were outlined and outcomes are typically a sequence of interesting qualities expressed vectorially [81].
Figure 7 Types of ML methods [81]
Figure 8. Supervised Learning in ML [81]
Conclusion

Machine learning, as a discipline of AI and one of the most popular categories of analytic techniques, is an important way for computers to learn. ML applications in materials science include novel material discovery, material property prediction, and other objectives spanning from the macro to micro level. The purposes of ML application in materials science are employed for quite diverse materials. A wide range of related studies shows that ML can be utilized to create precise and effective solutions for materials science. The areas in which ML can be used in materials science are becoming wider as theories and methodologies evolve.

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