HG-CAD: Hierarchical Graph Learning for Material Prediction and Recommendation in Computer-Aided Design

To support intelligent computer-aided design (CAD), we introduce a machine learning architecture, namely HG-CAD, that recommends assembly body material through joint learning of body- and assembly-level features using a hierarchical graph representation. Specifically, we formulate the material prediction and recommendation process as a node-level classification task over a novel hierarchical graph representation of CAD models, with a low-level graph capturing the body geometry, a high-level graph representing the assembly topology, and a batch-level mask randomization enabling contextual awareness. This enables our network to aggregate geometric and topological features from both the body and assembly levels, leading to competitive performance. Qualitative and quantitative evaluation of the proposed architecture on the Fusion 360 Gallery Assembly Dataset demonstrates the feasibility of our approach, outperforming selected computer vision and human baselines while showing promise in application scenarios. The proposed HG-CAD architecture that unifies the processing, encoding, and joint learning of multimodal CAD features indicates the potential to serve as a recommendation system for design automation and a baseline for future work. [DOI: 10.1115/1.4063226]

Keywords: hierarchical graph learning, graph neural network, computer-aided design, material recommendation, artificial intelligence, graphics processing unit (GPU), computing for design and manufacturing
1 Introduction

As a critical aspect in design automation and mechanical engineering, appropriate material selection is a demanding task that requires a devotion of time and expertise through joint analysis of performance, manufacturability, and sustainability [1,2]. Given this challenge, the integration of intelligent tools, such as the application of emerging machine learning algorithms to existing engineering systems, could be explored to assist designers with varying expertise by providing material selection recommendations learned from prior designs in an automated manner [3]. For example, intelligent agents trained on knowledge graphs generated from large-scale design data could help facilitate material selection processes that require extensive evaluation of trade-offs between diverse material candidates. However, there is an opportunity to expand on prior work toward later phases of the design process by extracting design knowledge from computer-aided design (CAD) models to complement existing semantic networks.

In the design domain, CAD tools are used to digitally create 3D models of physical objects and represent various design aspects, including the geometries, topologies, dimensions, tolerances, degrees-of-freedom, material information, and relative motions of components [4]. CAD tools consider material properties and interactivity characteristics between different materials, typically used for simulation to optimize the design or for rendering workflows. Both activities help designers assess and visualize trade-offs between different materials and identify the best materials for the design. Most recently, several large datasets of curated CAD models have been made publicly accessible that support machine learning methods for various data-driven design applications [5–7]. For creating representations of CAD models compatible with machine learning methods, graph data structures have been leveraged to represent design data and to capture various relationships, including semantic relationships between engineering and design concepts [8], joint relationships between parts in assemblies [9,10], and relationships between entities in boundary-representation (BREPs) [10–12]. Simultaneously in product design, efforts to consolidate design knowledge in graph-based knowledge structures have resulted in robust graph representations of domain-specific semantic relationships, which have proven useful for concept generation and evaluation [13–15].

In this paper, we present HG-CAD (Fig. 1), a learning-based approach to predict the material of each part in the assembly. Despite the abundance of 2D drawings and 3D CAD design repositories, the automated prediction of part materials in mechanical design remains challenging due to the ambiguity between the multi-modal design knowledge and their relationships to material selection. As illustrated in Fig. 2, the geometry and ground-truth materials of bodies might not have a one-to-one mapping. For example, a screw can be either ferrous or non-ferrous metal, depending on various design requirements such as cost, weight, corrosion resistance, adjacent parts’ material, the assembly it belongs to, and its function. This dependency on context introduces additional complexity for existing classification methods that rely on visual recognition. Therefore, their structural and contextual information must be considered to facilitate the accurate classification of bodies within assemblies. Leveraging the expressive power of graphs, we propose a graph neural network (GNN)-based architecture in which material prediction is posed as a node prediction task. Motivated by the importance of material selection to support design automation, this unified architecture is expected to aid designers in selecting appropriate materials by providing part-level material suggestions given a product assembly. We validate the method through quantitative and qualitative evaluation against computer vision and human baselines using the Fusion 360 Gallery Assembly Dataset [9].

To summarize, our work incorporates the following stages and contributions to the areas of design automation and engineering:

1. We study the material prediction task for design automation and devise a novel hierarchical graph representation of CAD assemblies to capture geometric and topological information from body and assembly levels.

2. We propose HG-CAD, which leverages the hierarchical graph representation of CAD assemblies for automatic prediction and recommendation of materials in unseen assemblies using graph neural networks while preserving contextual awareness using batch-level mask randomization.

3. We evaluate the effectiveness of the proposed architecture by conducting quantitative and qualitative experiments, demonstrating competitive performance as compared to
state-of-the-art learning-based models while providing insights and potential applications regarding strengths and limitations through a comparison with a human baseline.

(4) We provide open-source code and documentation for the proposed HG-CAD architecture (with data extraction, processing, baseline model variations, hierarchical graph construction, and learning toolkit)\(^2\) for reproducibility and further research while being fully compatible with the scalable Fusion 360 Gallery Assembly Dataset [9].

2 Related Work

We review prior work on material selection, machine learning algorithms based on deep neural networks, and representation methods tailored to CAD design automation. Reflecting on these past work, we provide deeper insights into the vision and motivation of the proposed work.

2.1 Material Selection in Product Design. In product design, material selection can be broken down into a general five-step procedure: (1) establishing design requirements, (2) screening materials, (3) ranking materials, (4) researching material candidates, and (5) applying constraints to the selection process [2]. Performance indices and material property charts, called Ashby diagrams, are often used to visualize, filter, and cluster materials [2,16]. Commercial tools such as Granta CES Edupack allow designers to compare thousands of materials based on user-input design requirements and constraints [17]. Product designers aim to meet customer needs and technical requirements, but the wide range of consumer products, manufacturing methods, and supply chain availability complicate the material selection process [18]. Accordingly, material aspects such as quality, cost, and function must be considered during product design as they directly contribute to the product’s success and its economic, environmental, and social impacts [19–21]. Material selection is further complicated when working with complex assemblies, as the material of individual parts also affects the assembly integration process [22].

Prior work on automated material selection focuses on specific classes of objects (e.g., nozzles and beams) [23–26] and specific design functions (e.g., heat transfer and storage) [27] and is often treated as an optimization problem where the best material is selected based on some required performance criteria [21,28,29]. Despite a few approaches being proposed to leverage neural networks (NNs) for material screening, they are limited in that they do not rank the selected materials [30]. Zhou et al. combine a two-layer NN with a genetic algorithm for selecting appropriate sustainable materials and validate their model on the design of a drink container [31]. Chandrasekhar et al. [26] leverage a variational autoencoder to project a discrete material database onto a differentiable latent space and couple it with a geometry encoder NN to simultaneously optimize the geometry and the material of a beam structure. Our method differs in that it only requires design information commonly documented in CAD, is agnostic to the manufacturing method and class of objects, and does not aim to find the most optimal material but suggests top-ranked appropriate materials to

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\(^2\)https://github.com/BrandonBian/hg-cad
the designer during the design process, thereby supporting user creativity and design flexibility.

2.2 Convolutional and Graph Neural Networks. Convolutional neural networks (CNNs) are a class of artificial neural networks predominantly used for feature representation and learning. Specifically, CNNs apply filters via sliding a kernel over the multi-dimensional input embeddings to perform convolution and pooling processes that extract, summarize, and represent the presence of detected features, which can be subsequently passed through a feed-forward neural network for representation learning tailored to tasks such as classification through activation. Due to the flexibility of the kernel dimensions, CNNs can effectively extract features of various scales of details that can be pooled into a unified representation and are thus most dominantly used for visual representation in computer vision [32,33].

Like CNNs, GNNs are a category of deep neural networks designed to perform learning and inference on data described by graphs and distributed in a non-Euclidean space. Message passing for representation learning in GNNs mainly consists of two main steps: aggregation, in which we collect features from topological neighboring nodes and edges; combination, in which we integrate the aggregated features into a single-node representation, later pooled to form graph representations. Specifically, GNNs learn representations over order-invariant data structured as graphs and often of variable sizes through an iterative process of transferring, transforming, and aggregating the representations with topological awareness [34–39]. The learned representations are combined into a graph-level representation, which can be used to perform tasks such as node, edge, and graph-level classifications.

In the proposed work, we leverage both types of neural networks to enable adequate representation and learning of CAD through joint exploitation of geometrical and topological features. In summary, the proposed architecture represents both CAD assemblies and bodies as hierarchical graphs, where the bodies are structured as graphs within assembly graphs and the assemblies are structured as super-nodes consisting of sets of body graphs, similar to the architecture proposed by Xing et al. [40]. CNNs are utilized for body-level geometrical representation extraction and encoding, whereas GNNs are adopted to perform topological representation learning with structural and contextual feature awareness. The proposed work’s combination of both neural networks in an end-to-end paradigm allows for joint geometrical and topological learning, thus exploiting the utilization of multi-modal CAD features to the full extent, allowing for classification and predictions with effectiveness and efficiency exceeding that of mono-modal representation learning.

2.3 Graph Representation of Computed-Aided Design Models. The design process is iterative and generates large amounts of data that can be organized and parsed for additional information that may be used to improve the design [41]. This information, collected from all aspects of the product life-cycle, is multi-modal and can be in the form of semantic names, customer requirements, 3D geometry, material properties, manufacturing tolerances, cost information, etc. These data may then be used to modify the design itself, enabling some design process automation by learning from prior examples [42,43]. Prior work looked at organizing and learning from design knowledge acquired from sources such as taxonomy-based design repositories [44,45], product tear-downs [46], patent data [8], and geometry-based design repositories [47–49]. When working with geometric data, graphs have been leveraged to represent BREP geometries with goals ranging from representing complex relations to solving design problems [11,50–52]. In particular, UV-Net [12] is a novel neural network architecture that can be leveraged for classification and segmentation tasks on B-rep data from 3D CAD models. UV-Net represents each assembly body as a face-adjacency graph, with face and edge features represented as a structured grid of UV-grid and U-grid of points, respectively.

Convolutional neural networks are applied on the face and edge features that are further processed through a graph neural network. The highly expressive nature of graphs in discrete encoding can also be applied to enable the more sophisticated representation and learning of multi-modal data. For example, Jones et al. [53] proposed a structured BREP graph convolution network that utilizes structured graph representation to encode heterogeneous BREP information, which prompted the creation of an assembly modeling tool for automatic mating of assemblies by effectively capturing the topological relations of parts. Similarly, Pfaff et al. [54] applied graph neural networks to mesh-based simulation, with an encoder that transforms mesh into graphs with the addition of edges, a processor that performs convolutional message passing, and a decoder that extracts signal for updates.

Our proposed work expands on prior literature by augmenting CAD representations through a novel hierarchical graph representation and learning architecture, effectively capturing the geometrical and topological features on both body and assembly levels while preserving contextual and structural awareness through GNN learning with node masking for CAD material prediction and recommendation.

3 Methodology

This section introduces the proposed HG-CAD architecture and the motivation behind the preprocessing steps for the CAD dataset. Next, we formulate the proposed hierarchical graph representation of CAD models and the detailed methodology for classifying and predicting assembly body materials using joint geometrical and topological learning with batch-level mask randomization.

3.1 Architecture Overview. As illustrated in Fig. 1, the proposed material prediction network consists of two primary branches: (1) the body-level module that extracts and processes the CAD body features, represents them in terms of body graphs to capture topology, and encodes the geometrical and physical features using different tools and transformations; (2) the assembly-level module that provides a graph representation of the entire assembly, passes the topological information and the body-level features collected from the previous module through a graph neural network for embedding generation, and using a densely connected neural network for representation learning and prediction.

3.2 Dataset and Preprocessing. The proposed architecture is compatible with the publicly available Fusion 360 Gallery Assembly Dataset [9]. As illustrated in Fig. 3, each assembly in the dataset has assembly-level feature properties, which are shared across numerous bodies connected via assembly relationships. Each assembly body has a set of body-level design features, and the bodies are manually organized by users in a hierarchy of occurrences, which are the building blocks that make up the assemblies. The dataset contains 154,468 bodies that are grouped into 8251 assemblies, created across different industries and design categories and with various levels of detail. This dataset was selected because of its scalable size, abundant diversity of designs and multi-modal features, ease of processing, and presence of body-level material labels. Due to the complexity and diversity of the dataset, preprocessing and feature encoding steps are performed to target the assemblies and bodies of interest and encode the extracted multi-modal raw features into comprehensible formats in the learning architecture. The preprocessing steps on the dataset are as follows.

Data filtering. We drop assemblies that are entirely labeled with the default material (Steel), resulting in 6336 assemblies.

Material label transformation. Each assembly body within the assembly dataset is defined with two material labels: a physical material label that defines the physical and mechanical properties
of the assembly body for simulation and an appearance material label used for displaying and rendering the object. Despite the different labels serving distinct purposes, it may be assumed that the designer intends to have each assembly body defined by one material. To reflect this, a transformation process is performed in which the non-default appearance label is selected as the material ground truth to replace the default physical material label, if applicable. This is performed to improve the quality and abundance of ground-truth labels while attempting to preserve the original design intent.

Material label grouping and dropping. Due to the diversity, skewness, and sparsity in the distribution of user-defined body-level materials, a manual regrouping step is taken to reorganize the ground-truth labels. Specifically, the detailed material labels, as transformed from the previous step, are mapped to their corresponding generalized material categories as defined in the Autodesk Fusion 360 material library, which are then regrouped into eight simplified material groups, as summarized in Table 1, based on a combination of their primary, secondary, and tertiary subcategories. After performing the regrouping of material categories for body-level ground-truth labels, the following two groups are dropped: Metal_Ferrous_Steel, which contains only the default carbon steel material and is thus less relevant to the material prediction task, and Paint, which contains generic and rendering material that is ambiguous and does not reflect meaningful design or physical properties. More specifically, the motivation for dropping the Metal_Ferrous_Steel label is double-fold: (1) it is configured as the default material label for design bodies in the Autodesk Fusion 360 platform, which makes it irrelevant as they do not contain sufficient user-configured information to benefit the model’s training, and (2) it is a dominant label that introduces label imbalance which induces prediction bias. Therefore, the regrouping step of ground-truth labels effectively reduces the complexity and distributional skewness while preserving logical correctness, whereas the dropping of default and paint-related material groups alleviates the confusion incurred during the architecture learning process. In addition, incorporating the final six simplified material groups in the prediction pool enables more flexible and scalable recommendation outputs than solely considering the exact material labels, thus leaving room for the creativity and freedom of designers.

### Table 1 Table of the simplified ground-truth material groups

<table>
<thead>
<tr>
<th>Simplified material group</th>
<th>Definition</th>
<th>Example(s)</th>
<th>Label count</th>
</tr>
</thead>
<tbody>
<tr>
<td>Metal_Aluminum</td>
<td>Aluminum-based metal</td>
<td>Aluminum alloy</td>
<td>10,606</td>
</tr>
<tr>
<td>Metal_Ferrous</td>
<td>Ferrous metal (excl. carbon steel)</td>
<td>Cast iron</td>
<td>7138</td>
</tr>
<tr>
<td>Metal_Ferrous_Steel (dropped)</td>
<td>Carbon steel</td>
<td>Carbon steel</td>
<td>39,891</td>
</tr>
<tr>
<td>Metal_Non-Ferrous</td>
<td>Non-ferrous metal</td>
<td>Platinum, silver</td>
<td>16,276</td>
</tr>
<tr>
<td>Other</td>
<td>Uncategorized material</td>
<td>Glass, fabric</td>
<td>15,028</td>
</tr>
<tr>
<td>Paint (dropped)</td>
<td>Generic, rendering, and coating material</td>
<td>Metal flake</td>
<td>13,193</td>
</tr>
<tr>
<td>Plastic</td>
<td>Plastic</td>
<td>Thermoplastic</td>
<td>20,063</td>
</tr>
<tr>
<td>Wood</td>
<td>Natural and engineered wood</td>
<td>Softwood</td>
<td>9107</td>
</tr>
</tbody>
</table>

3.3 Hierarchical Graph Representation. Despite multimodal expressive features, the CAD assemblies and their corresponding bodies are difficult to encode and may introduce complications when directly delivered into machine learning architectures. Therefore, we represent CAD assemblies using a graph representation, wherein, body attributes are captured as node features and topology is captured using edges:

#### 3.3.1 Body-Level Graph Representation. As illustrated in Fig. 4(c), we define body graphs as attributed and directed face-adjacency graphs representing CAD assembly bodies, where the visible parametric surfaces are represented as nodes, and the visible interval of the parametric boundary of faces are represented as edges that connect two faces that are adjacent to each other. For graph feature generation, we follow the methodology as introduced in UV-Net. Specifically, we generate the body graph node features by parameterizing assembly body surfaces as sets of 2D features constituted of the absolute 3D normalized coordinates sampled from surface domains with a uniform step size. Similarly, body graph edge features are created by parameterizing boundaries as sets of 1D features constituted of absolute point coordinates sampled from its parameter domain with a uniform step size. The node and edge feature matrices generated by a stacking of individual feature vectors, along with the connectivity information of nodes, are organized into deep graph library (DGL) [55] graph object instances and are subsequently delivered into the learning
architecture along with the assembly-level graphs created according to the subsequent section.

3.3.2 Assembly-Level Graph Representation. As illustrated in Figs. 4(a) and 4(b), we define assembly graphs as attributed and directed multi-graphs representing entire CAD assemblies, where bodies corresponding to individual parts are encoded as graph nodes. Connections representing assembly relationships between the bodies are encoded as edges between corresponding nodes.

When constructing assembly graph node features, we consider the following three assembly body physical properties: body area represented in square meters, body volume represented in cubic meters, and body center of mass represented in the x, y, z coordinates. All physical properties are generated by Autodesk Fusion 360 and are numerical. Standard scaling transformation is applied to each physical property to generate their corresponding physical feature embedding vectors. We further consider the features defined at the assembly level and are thus shared among the bodies of the same assembly globally. These features include the assembly physical properties and assembly geometric properties, which are integrated from individual assembly body properties, as well as design category, industry, and products that the assembly belongs to. Generally, global features represented as floating-point or integer numbers are preserved and normalized using standard scalar transformation, whereas categorical features represented as character strings are one-hot encoded. When constructing assembly graph edge features, we obtain the following three predominant types of connection relationships: contacts define the relationship between two bodies whose faces are in contact with each other; joints define the relationship between two bodies whose relative pose and degrees-of-freedom are constrained; occurrence relationship represents the user-assigned relative hierarchy of the bodies, such as multiple bodies sharing the same occurrence, or distributed across a parent–child occurrence relationship. These three connection types, being categorical, are encoded using the one-hot method and serve as the edge features.

The resulting graph is attributed since each node and edge is attached with features that effectively capture the physical and structural information. The graph is directed since each edge contains a source node, a destination node, and a direction that correlates the two nodes. Furthermore, the graph is a multi-graph since it allows the existence of multiple edges between any pair of nodes. Once the assembly graphs are constructed, instances with less than three nodes or less than two edges are considered trivial and are thus discarded. Following the reasoning in Sec. 3.2, the nodes with ground-truth material category labels of Metal_Ferrous_Steel and Paint are dropped, and the edges connected to them are removed accordingly from the connectivity matrices. The node and edge feature matrices are generated by stacking individual feature vectors and the shared global features. The feature matrices and the connectivity information of nodes represented in coordinate (COO) format are organized into PyTorch Geometric [56] graph objects and are subsequently delivered into the learning architecture along with the body-level graphs created according to the prior section.

3.4 Hierarchical Learning Architecture. Building on the hierarchical graph representation of CAD assemblies, we present the learning architecture of HG-CAD, in which we utilize graph neural networks to generate graph embeddings in both body and assembly hierarchies by passing node-level feature messages through convolution.

We denote a set of assembly graphs of interest constructed from Sec. 3.3.2 as $G = \{G_1, \ldots, G_{\text{assembly level}}\}$, with individual graphs $G_i = (V_i, E_i, Y_i, M_i, \varphi_i)$, each of which consists of a set of body nodes $V_i = \{v_1, v_2, \ldots, v_{|V_i|}\}$ (carrying node features $h_i \in \mathbb{R}^{d_j}$), a set of assembly relationship edges $E_i = \{e_{l, j} = (v_l, v_j)\} \subseteq V_i \times V_i$ (carrying edge features $h_{l, j} \in \mathbb{R}^{d_e}$), a set of body graph embeddings $G_i = \{g_1, g_2, \ldots, g_{|V_i|}\}$ corresponding to each body node, a masking matrix $M_i = \{m_1, m_2, \ldots, m_{|V_i|}\}$ such that $m_i \in \{0, 1\} (i \in \{1, \ldots, |V_i|\}$) representing the validity of body graph embedding, and a set of ground-truth material category labels $Y_i = \{y_1, y_2, \ldots, y_{|V_i|}\}$ corresponding to each node. The topological representation learning process can be formulated as a supervised node classification task, in which a node-level representation $h_i (V_i \in V)$ is learned from a combination of the assembly graph topology, assembly graph node and edge features, and the body-level geometric embedding, such that the ground-truth material category label for each body can be predicted.

During the training time, at a certain $l \in \{1, \ldots, L\}$ layer of the graph neural network, message passing is performed between direct neighbors through neighborhood aggregation, where the representation of a certain node $v_i \in V_i$ is iteratively updated by a combination of aggregated neighboring node and edge features, as defined below:

$$h^{(l)}_i = \phi_i \left( h^{(l-1)}_i, \bigcup_{v_j \in N_i} \phi_j \left( h^{(l-1)}_j, h^{(l-1)}_{\text{edge}} \right) \right)$$

where $h^{(l)}_i$ denotes the representation of node $v_i$ at layer $l$ of the network, and $\phi(.)$ and $\phi(.)$ denote the parametric combination and aggregation functions, $N_i$ denotes the topological neighbors of node $v_i$, and the representation of node $v_i$ during the aggregating step contains a concatenated feature of body-level geometric representation generated from the DGL body graphs $g_i$ and the corresponding masking matrix for the elimination of invalid
embeddings, as defined below:

\[ N_i = \{ v_i \in V : \exists e_{ij} \in E \}, \quad h_i = [h_i^g, g_i^f, m_i] \]  

(2)

where the body graph geometrical embedding of a graph node at a given layer \( g_i \) is generated by a summation aggregation of the body graph’s node features (2D convoluted from parameterized surface coordinates) and edge features (1D convoluted from parameterized boundary coordinates) using a multi-layer perceptron, following the learning architecture as introduced in UV-Net [12].

GraphSAGE [57] is selected for the assembly-level graph neural network layers, utilizing a mean aggregator. The graph connectivity information, concatenated node, and edge feature matrices are passed into the GNN encoder during the training process. The node and edge embedding is obtained after message passing of each neural network layer, and non-linear activation via leaky rectified linear unit [58] is performed. For generating the final node embedding to perform node classifications on the input graph, a similar procedure as the jumping knowledge networks [59] is performed by aggregating the node embedding of individual GNN layers from the learning architecture through max-pooling. The final embedding is then passed through a multi-layer perceptron with batch normalization layers [60], parametric rectified linear unit [61] for non-linearity activation, and softmax activation for predicting class distributions. Weighted cross-entropy loss is adopted to prevent the neural network from overlooking rare classes. The weights for each class are initialized as inversely proportional to the ground-truth class frequencies.

Considering application scenarios, we propose batch-level mask randomization, in which ground-truth material category labels are injected into portions of assembly graph nodes using randomized masking to prompt the neural network learning toward fully exploiting the contextual design knowledge and user-introduced information. Motivated by the fact that designers may have access to material information for completed portions of their design assemblies, we inject ground-truth material category labels as additional material information for completed portions of assembly graph nodes using randomized masking to prompt the neural network learning toward fully exploiting the contextual design knowledge and user-introduced information. In the case of the proposed model and the baselines to ensure reproducibility, as summarized in Table 2 and detailed in the following paragraphs.

### 4 Experiments

In this section, we analyze and evaluate the performance of our architecture through experiments on the Fusion 360 Gallery assembly dataset. Specifically, we consider the following three baselines of machine learning classification models: (1) UV-Net [12], which exploits geometry as sampled from solid surfaces and edges together with topology structured as graphs; (2) PointNet [62], which samples point clouds on mesh surfaces to perform object classification on CAD models; (3) MVCNN [63], which uses a CNN architecture that combines 2D geometry obtained from multiple views of a 3D model to perform recognition and inference. We also consider a human baseline, for which we ask students from mechanical engineering backgrounds to provide manual body-level predictions on existing design assemblies.

#### 4.1 General Setup

To ensure a fair comparison, we split the constructed assembly graphs obtained from Sec. 3.3.2 into the training, validation, and test sets, which remain fixed across all models (but excluding the human baseline). Specifically, we generated the training, validation, and test sets based on the proportions of 64%, 16%, and 20%, respectively, through a random sampling process on the assembly graphs, repeated for a total of three iterations with different random seeds. Furthermore, we provide the configuration of the proposed model and the baselines to ensure reproducibility, as summarized in Table 2 and detailed in the following paragraphs.

**UV-Net.** By removing from our model the assembly-level GNN, we are left with only the part-level GNN, which leverages UV-Net to represent the part geometry. We use the official implementation of the UV-Net [12] classification model where the final linear layer was modified to output six classes. Each B-rep body is converted into face-adjacency graphs with 2D and 1D UV-grids as node and edge features, respectively, where we used ten sample points for the u- and v-directions as in the original implementation. The model was trained with the Adam optimizer [64] for 350 epochs, and the weights of the model corresponding to the minimum validation loss were used in our experiments.

**PointNet.** For each input body-level geometry encoded as triangulated B-Rep representation, 2048 points are randomly sampled. An Adam optimizer with a learning rate of 0.001 and a momentum of 0.9 is used. Sparse categorical cross-entropy is used to calculate loss, and a dropout rate of 0.3 is used on the last fully connected layer of hidden dimension 256 to alleviate overfitting. PointNet is implemented using TensorFlow, and the model is trained until the convergence of validation loss.

**MVCNN.** The MVCNN is trained using a PyTorch implementation and uses the ResNet architecture [66] with a supervised learning regime. A patience factor is used to stop the training process

![Table 2](https://github.com/AutodeskAILab/UV-Net)
after 20 epochs, which increases the validation accuracy, resulting in around 30 training epochs. The models are trained with eight as the batch size, 512 embedding dimensions, 12 views, and $1 \times 10^{-4}$ learning rate.

**Human baseline.** To compare the performance of the proposed work to that of humans, we asked three Master’s students to provide their predictions of material category labels for assembly bodies. Specifically, we randomly sampled 300 assemblies from the entire dataset and dropped the assembly bodies whose ground-truth material groups were *Metal_Ferrous_Steel or Paint*, simulating the process as described in Sec. 3.2. To ensure fairness of comparison, we sought to reproduce the information provided to the baseline models by presenting the human labelers with an interactive user interface containing the following context: 2D thumbnail images of the assembly body and its corresponding assembly, 3D rendered and interactive display of the assembly geometry with the body portion highlighted, semantic names and physical properties of the body, as well as the assembly-level global features.

For each assembly body, the students are asked to classify its material into one of the six material groups listed in Table 1 based on their experience, expertise, and the provided information. Figure 5 shows an example human labeling template.

### 4.2 Quantitative Evaluation.

Table 3 summarizes the quantitative results of the experiments compared between the proposed model and the baselines. To evaluate the multi-class classification performance and consider the skewness of ground-truth material group distribution, we record the micro $F_1$ score calculated by weighting each prediction instance equally. The performance statistics of each model are recorded per iteration and are averaged across the iterations to calculate the mean and standard deviations. For evaluating the human baseline, the human labelers’ body-level predictions are concatenated and compared via one-to-one mapping to the corresponding assembly body’s ground-truth material group. Due to having only one iteration over the 300 assigned assemblies, the human baseline performance does not have a standard deviation. From the numerical results, we infer that material prediction in CAD is generally a highly challenging task for both machine learning models and human labelers, due to the highly variational distribution of body geometry and assembly ground truth. The proposed HG-CAD, when producing predictions of all body materials of an assembly without referencing contextual material information, produced results that are only sparingly above that of the baseline. However, when enabling the joint learning of assembly topology and part geometry with contextual information using node masking, the proposed architecture achieves an average of 0.59 micro $F_1$ score, which surpasses that of the baseline machine learning models that mainly rely on geometry. This supports our hypothesis that the task of material selection depends on multiple factors other than the geometric features of the design, namely the topology of assembly bodies that enable contextual awareness during hierarchical graph learning. To further investigate the performance discrepancies and the human baseline results, we plot the classification confusion matrices for the models displayed in Fig. 6.

![Fig. 5](image)

An ideal confusion matrix for a classification model with the best classification should demonstrate clear diagonal patterns with no shades in other regions. From the confusion matrices, we observe that UV-Net and PointNet misclassify most material categories as Plastic, as observed by the significant shades in the plastic column. This is because the Plastic category is a majority class, with many samples dominating the other classes. The imbalance of label count could have misguided these two models to make predictions biased toward the Plastic category. UV-Net demonstrated less confusion and thus better classification performance than PointNet, possibly due to the inclusion of body-level topology with graph representations. MVCNN showed a clearer diagonal pattern, but confusion still persists with the Metal_Non-Ferrous and Other groups. The proposed model, deprived of material context information shared during message passing and learned via node masking,

<table>
<thead>
<tr>
<th>Table 3</th>
<th>Table of the experimental results for the material prediction task</th>
</tr>
</thead>
<tbody>
<tr>
<td>HG-CAD (no mask)</td>
<td>UV-Net</td>
</tr>
<tr>
<td>Acc.</td>
<td>0.59 ± 0.031</td>
</tr>
</tbody>
</table>

**Fig. 5** An example graphical user interface display for human baseline.

Transactions of the ASME
demonstrated similar confusion as the computer vision baselines, though with a slight alleviation in severity. Significant confusion is observed for the human baseline, where no assembly body was classified as Metal_Aluminum. This may be because Metal_Aluminum is a precise and exclusive class compared to the other five more general and inclusive categories, and human labelers confused aluminum material with a non-ferrous metal material. Compared to the four baselines, HG-CAD’s classification performance is superior, showing a much clearer diagonal pattern and, thus, less confusion during prediction. There is still a slight prediction pattern biased toward the majority class of plastic, but the effect is minimal. These results, together with the observation illustrated in Fig. 2, demonstrate that representation learning on geometry only is insufficient for making body-level material prediction and classification and that the inclusion of topology via graphs as well as neighborhood context via random masking can greatly reduce confusion.

4.3 Qualitative Evaluation. To investigate the prediction results in a real-world use case setting and to gain further insights for potential application, we provide a set of rendered assemblies to compare the prediction results of our model and that of the baselines in a qualitative manner. Specifically, we select four assemblies from a union of the test set of the machine learning-based models and that of the human baseline. The chosen assemblies are delivered into the machine learning models pre-trained on the fixed train-test split, and the inference results of body-level material predictions are produced. Subsequently, the 3D models of the chosen assemblies are rendered with colors corresponding to the predicted material categories.

Illustrated in Fig. 7 is the visualized qualitative comparison. The gray color represents the material categories Metal_Ferrous_Steel and Paint, which are dropped and therefore do not participate in the evaluation process. From the visualized comparison, our proposed model can function well when dealing with bodies of similar geometry and symmetrical distribution in space. For example, our model predicted almost perfectly the material category of the pistons in the first sample assembly, whereas the predictions of the other baseline models are rather arbitrary. This demonstrates that our model can effectively exploit the body-level geometrical information in conjunction with assembly-level topological information through the proposed joint representation learning methodology. For the performance on the latter three assemblies, the baseline models gave predictions inconsistent with the ground truth, often confusing the aluminum metal category (blue) with the non-ferrous metal category (orange). A similar confusion within the metal categories can be observed for the human baseline predictions, where almost all metal occasions are misclassified as Metal_Ferrous. The qualitative evaluation demonstrates the effectiveness of the proposed joint learning of topological and geometrical representation methodology compared to baseline methods that rely on only a single aspect of CAD representation.

5 Discussion

5.1 Limitations and Improvements

5.1.1 Data and Model Augmentation. While running single-node prediction experiments, we observed that the inference confusion decreases with increasing training and validation assemblies. Despite the batch-level randomized node masking process, only one node per assembly graph participates in the loss calculation for training and validation; thus, the support for each class is diminished, which may lead to problems such as overfitting. Therefore, additional steps should be taken to perform data and graph augmentation to alleviate this issue. For collecting additional CAD data with a more varied distribution of features and ground-truth materials, we seek to expand the currently adopted Fusion 360 Gallery Assembly Dataset by inviting more designers to contribute or by using additional publicly available datasets such as ABC and Auto-Mate [6,53]. For graph-level augmentation, existing graphs can be manipulated to generate additional training and validation samples, such as by creating sub-graphs or synthesized graphs from past designs through node dropping, edge dropping, or sub-graph sampling [67]. Existing tools designed to support automated reasoning, such as Open-NARS [68], could also be applied to the machine learning process to support the inference and imputation of rare material classes. Since neural networks require abundant high-quality training samples, incorporating the reasoning system might provide a more balanced approach through deduction.
abduction, and induction reasoning. In addition, future work can be directed to experimenting with a more diverse set and combination of the neural network building layers. In the current iteration, we examined the performance with graph attention networks (GATs), graph isomorphism networks (GINs) as well as linear transformation layers as suggested in Ref. [31], and observed significant overfitting of the training dataset due to the uneven distribution of assembly architectures and material selections, as indicated in the convergence graph of Fig. 8. Furthermore, due to the highly variable discrepancies between the topology and structure of design assemblies, more experimental iterations should be performed with diverse splits. Similarly, we also intend to extend our human baseline sample size by incorporating human participants from different realms of engineering and expertise levels for a more comprehensive investigation and analysis of human engineers’ prediction patterns.

5.1.2 Functional and Behavioral Information. The current stage of work focuses on the structural aspect of objects, including the geometric and physical properties of bodies and the assembly relationships that correlate them. While structural analysis has proven to be successful in the derivation and tabulation of material performance indices for standard mechanical design cases [69], the functional and behavioral aspects of the design are also highly influential to material selection [2,70]. Therefore, one promising improvement would be incorporating the functional information (i.e., the purpose) and the behavioral information (i.e., the attributes) of bodies as node features to provide more context for learning. Functional and behavioral information might be inferred from the name of parts in the assemblies, and prior work has shown promise in learning from these natural language labels commonly found in CAD [44,71], and large language models could be leveraged for transfer learning. While body-level semantic names might implicitly represent some functional and behavioral information, introducing additional user inputs (e.g., cost and size requirements) would enable predictions tailored to specific design requirements.

5.2 Future Plans

5.2.1 Regression Model for Material Properties. One limitation of the proposed work is its dependency on the Autodesk Fusion 360 Assembly Dataset and its categorical material library. Specifically, new designs from the Fusion 360 Gallery can be imported as additional data during training and inference. For a new assembly design acquired from the Autodesk Fusion 360 gallery, it can be encoded as a JSON file automatically by the FUSION 360 software along with the thumbnail images and 3D object files, where the embedding dimensions of the body-level features are automatically handled by the underlying UV-Net architecture. The dimensions of the graph embedding are determined by the hyperparameters defined by the user in the HG-CAD architecture. Despite alleviating possible confusion by dropping bodies of default and paint material, the regrouping process may introduce bias that limits the trained model’s application to only materials found in the original dataset, thus limiting scalability and might not suit the needs of designers in different industries. A possible improvement might be to restructure the problem as a regression task and develop a model to predict relevant physical and mechanical properties involved in the selection of materials, expanding prior work that trained NNs directly on material properties [26,72]. Moreover, by mapping relevant material properties of the material library of the training data onto an Ashby chart, clustering the material properties on the chart may enable a more flexible material selection method.

5.2.2 Graph Predictions and Similarity Search on Computer-Aided Design Models. We envision the potential expansion of our work in producing diverse predictions for CAD automation that are not limited to material selection. Specifically, we seek to incorporate graph edge prediction and global context prediction functionalities by varying our current learning architecture. For graph edge prediction task, we aim to support the user-design process of organizing and correlating assembly bodies by providing insights into their hierarchical and relational information through

<table>
<thead>
<tr>
<th>Ground Truth</th>
<th>HG-CAD</th>
<th>UV-Net</th>
<th>PointNet</th>
<th>Human Baseline</th>
</tr>
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</table>

Legend:
- **Metal_Algum**: Blue
- **Metal_Ferrous**: Yellow
- **Metal_Non-Ferrous**: Red
- **Plastic**: Green
- **Wood**: Orange
- **Dropped**: Dark Blue
- **Other**: Light Pink

Fig. 7 Qualitative comparison of material prediction results for selected assemblies
representation learning based on their properties and that of the assembly. For global context prediction, we aim to provide users with an informative overview of the entire assembly design through representation learning tailored to the global graph features shared across individual bodies. Furthermore, the feature embedding of the entire assembly graph representation encoded with assembly-level topology and body-level geometrical information can be used with unsupervised learning methods, such as high-dimensional clustering, to enable similarity search on CAD models with structural and contextual awareness. One application scenario would be a post-design recommendation, in which we retrieve, based on similarity search results, past CAD designs that are similar to the newly produced user design, which may serve as sanity checks or baselines for the designer to reference. While the current solution only considered GraphSAGE, GAT [73], and GIN as the building layers of the graph encoder, we envision that additional GNN layer types provided by the DGL library, such as RelGraphConv and CFConv, can be incorporated in future work, provided the flexibility of the model architecture. Furthermore, we plan to seek further collaboration and a more diverse design repository access to apply our HG-CAD architecture. We believe that our proposed architecture is flexible and competent enough to handle a wide range of use cases and anticipate that it may be applicable to high-value components and assemblies, satisfying the needs of broader users.

5.2.3 A User-Guided Active Learning Process. We want to overcome the limitations of learning solely from existing designs and would want to preserve the space for creativity by prompting a user-guided learning process in future works. Specifically, we envision an active learning process for the single-node prediction task, in which the model not only provides a single-shot prediction for the user-input design based on its learned knowledge from existing designs but also actively adjusts its weights and parameters as the user introduces new ground-truth information to existing bodies or even adding new bodies to the design. Another possible future direction to bring the user into the learning loop is to allow post-prediction feedback and adjustments. In the current iteration of the project, we evaluated the efficacy of the human labelers by collecting their feedback on the challenges during manual prediction, their confidence in their results, and how they think the entire experimental procedure can be improved. From the collective
feedback, we found that the major challenge that our human subjects encountered during their predictive labeling process derives from a lack of global context, which confounds their judgement on bodies of similar geometry. We hope this piece of information can demonstrate how the labeling of assembly part information can be hard to evaluate, thus highlighting our work’s goal of automating the process by leveraging machine learning’s capability of capturing diverse and latent representations from large repositories of past data to generalize better than human subjects. To provide more information that can bring the user into the loop, after displaying the top predicted material categories along with their corresponding confidence scores, we prompt the user with several questions, such as the requirement for the target assembly body’s weight, volume, and cost, and adjust our predictions or re-run the inference step based on the newly introduced information. This should effectively narrow the search space and thus predict a scope tailored to the user’s specific needs.

6 Conclusion

In this paper, we propose HG-CAD, a unified architecture based on a hierarchical graph representation that enables intelligent computer-aided design by predicting the material category of bodies in design assemblies through joint learning of geometry and topology from both body-level and assembly-level scales in an end-to-end procedure, and with contextual awareness enabled through batch randomized node masking. Furthermore, we present a systematic workflow for the automated feature extraction, encoding, and graph representation generation of CAD models compatible with the predominant and scalable Fusion 360 Gallery Assembly Dataset.

We compare our proposed model with three state-of-the-art learning-based models and a human baseline for the experimental evaluation. Specifically, we formulate the body-level material prediction task as a node classification task on graphs, where we randomly generate the batch-level Boolean indicator masks to allow the network to make predictions on individual target nodes while referencing the ground-truth information from topologically neighboring nodes. Quantitative results demonstrated the effectiveness of the proposed methodology of representing CAD models as hierarchical graphs to enable joint learning of body-level geometry and assembly-level topology with contextual awareness in material prediction. Qualitative comparison using visualized predictions also reflected the potential of our proposed model in capturing and utilizing the similarity in geometry in conjunction with the topological symmetrical characteristics, showing promise in supporting human-in-the-loop design automation applications. Specifically, we believe that the proposed method’s capacity to accommodate large-scale databases and flexibility in incorporating the designer’s knowledge can be used to create a recommendation system for users by learning best practices from existing designs. Furthermore, the architecture may serve as a baseline or foundation for future works leveraging graph neural networks for design automation.

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Conflict of Interest

There are no conflicts of interest.

Data Availability Statement

The datasets generated and supporting the findings of this article are obtainable from the corresponding author upon reasonable request.

Nomenclature

AI = artificial intelligence
ASME = American Society of Mechanical Engineers
AIEDAM = artificial intelligence for engineering design, analysis, and manufacturing
CC = global clustering coefficient
CIE = computer in engineering
CPL = characteristic path length
CC-WS = Watts-Strogatz local clustering coefficient
CPLRG = characteristic path length of the equivalent random graph
DTM = design theory and methodology
DP = design parameter
FR = functional requirement
GED = graph edit distance
IFIP = International Federation for Information Processing
ME = mechanical engineering
SWI = small-world index
TRL = technology readiness levels

References