A Survey of Automatic Text Summarization using Graph Neural Networks

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Abstract

Although automatic text summarization (ATS) has been researched for several decades, the application of graph neural networks (GNNs) to this task started relatively recently. In this survey we provide an overview on the rapidly evolving approach of using GNNs for the task of automatic text summarization. In particular we provide detailed information on the functionality of GNNs in the context of ATS, and a comprehensive overview of models utilizing this approach.

1 Introduction

The advent of the internet has led to an explosion in the amount of textual information available online. The extensive availability of textual information paired with a need to quickly understand it has led to major efforts in the field of automatic text summarization (ATS). For a comprehensive review and survey of ATS as a task we recommend a recent survey by El-Kassas et al. (2021).

The goal of ATS is to produce a concise, correct and fluent summary of a given text. Although this definition is intuitively understandable, there is no commonly agreed upon formal definition for these qualities. This is in part due to the difficulty of the task itself as producing a summary with the above properties is challenging, even for humans.

Although the field of ATS has made major steps forward, generally, we differentiate between two basic approaches to the task of ATS. The first being extractive ATS and the second being abstractive ATS. Extractive ATS involves extracting text spans from the original document such that a summary is generated. Thus the main challenge consists in identifying useful spans of text from the original document. This approach is popular as it eliminates the non-trivial task of generating factually correct and coherent sentences. Abstractive ATS on the other hand involves the challenging task of generating novel sentences for a summary. Despite impressive advancements, generating factually correct and fluent sentences is still a major challenge in ATS (Kryściński et al., 2020). In addition to the above, one differentiates between the task of single-document and multi-document summarization.

1.1 Why graph neural networks?

Contemporary solutions to the task of ATS suffer from a number of issues, chiefly an inconsistent evaluation protocol and, somewhat, a lack of progress, as noted by Kryściński et al. (2019). In recent years GNNs have been successfully applied to a number of downstream NLP tasks such as classification (Liu et al., 2020b) (Zhang et al., 2020b) and translation (Xu et al., 2021) (Yin et al., 2020). Although GNNs may not be able to solve all problems related to the task of ATS, we believe that they can at least give a new perspective to this task. Generally GNNs bring a number of advantages to ATS which we believe to be significant enough to warrant further research, and this survey. In particular we want to highlight the following aspects of GNNs:

- **Scalability and Flexibility.** A vast number of ATS models are based on BERT (Devlin et al., 2019). However, the computational complexity of BERT-based ATS models grows quadratic with the input length; due to the self-attention operation. This fact renders them impractical for long, or even medium sized text documents. Recently some work has been done in order to circumvent this limiting factor (Ding et al., 2020) (Zhang et al., 2020b). In contrast, GNNs can scale by their nature to graphs of thousands of nodes and more. This is in part due to the linear scaling of the memory cost with regards to the input size. The total memory cost of a GNN model depends on the size of the graph, the number of layers and the feature vector size of the nodes present. Formally, for $L$ layers and an input
of $N$ nodes with each node’s feature vector being of size $H$ the memory complexity is $O(LNH)$. But even for very large graphs on the scale of millions of nodes one can utilize GNNs. This can be achieved using methods such as neighbour sampling or distributing the graph over multiple GPUs, as done for example by Jia et al. (2020b). We recommend the paper by Li et al. (2021) for insights as to how one can train large and very deep GNNs. As the input of a GNN is a graph, the input can vary in size, therefore GNNs are also able to cope with changing text sizes and structures. Both of these aspects combined allow GNNs to produce summaries which are not restricted by hard-coded limits related to input or output size.

- **Understanding and Explainability.** It is often difficult to understand why a model arrived at a certain conclusion. Additionally it is often difficult to see how the model aggregates information. This is not the case with GNNs, as with the help of methods such as GNN Explainer (Ying et al., 2019) one can understand which nodes were used by the model to reach its output. This removes a layer of the black-box magic present in many current non-GNN models. We recommend the survey by Yuan et al. (2020) for an overview of methods for generating explanations for GNNs.

1.2 Related Surveys

As the application of GNNs to ATS is rather novel, to the best of our knowledge there is only one survey on the topic. The survey by Luo et al. (2020) gives an introduction to the topic. However, it does not provide much detail on GNNs in the context of ATS, nor does it cover all models in the space with a taxonomy.

As for GNNs themselves, there exists a large number of surveys on GNNs as a technology. In particular, we want to highlight the surveys by Wu et al. (2020b) and Zhou et al. (2020) which provide a general overview on GNNs and their applications in different fields. The survey by Abadal et al. (2021) provides more technical and theoretical details on GNNs. The survey by Wu et al. (2021a) is of particular interest as it focuses on the usage of GNNs for NLP as a domain. Additionally, we want to note here the survey by Wu et al. (2020a) on the usage of GNNs in recommender systems and the review by Malekzadeh et al. (2021) on the usage of GNNs for text classification.

For a more theoretical approach we point the reader to the analysis by Xu et al. (2018) which establishes a number of important theoretical properties for GNNs. An analysis of the Vapnik–Chervonenkis (VC) dimension of GNNs was performed by Veličković et al. (2018).

In general, the contributions of our survey are as follows:

1. We provide a detailed explanation of GNNs in the context of ATS.
2. We introduce a simple taxonomy for GNN models used for ATS.
3. We provide a comprehensive overview of innovative GNN models for ATS and discuss further directions for future research.

The rest of survey is structured as follows. First we will give a comprehensive explanation of GNNs in the context of ATS. Next, we will explore a number of interesting and innovative models. Finally, we will finish with a conclusion and an outlook on the future usage of GNNs for ATS.

2 Graph Neural Networks

Graph based methods for ATS are not a new innovation, with methods such as TextRank (Mihalcea and Tarau, 2004) being first presented in the early 2000s. In fact, the core idea of early graph-based approaches such as TextRank is similar to the one used by GNNs designed for the ATS task. The idea is to encode the structural and semantic information contained within a text document into an explicit form with the help of a graph.

Deep learning has by now become a common tool for solving tasks throughout many domains, with various end-to-end paradigms such as recurrent-neural-networks (RNNs) or convolutional-neural-networks (CNNs) emerging as versatile and powerful tools. In particular deep learning has shown extraordinary power for data which is Euclidean in nature. As an example, images can be represented as regular grids in Euclidean space. Using such a representation CNNs are able to extract meaningful local representations with the help of convolution. However, for many domains and applications such a representation is often either inconvenient or not even directly possible. The most obvious example being the field of
chemistry in which molecules should be modeled as graphs. The complexity of translating such complex data into existing deep learning paradigms has led, in part, to the development of GNNs which attempt to leverage the power of deep learning for non-Euclidean data.

2.1 Defining the Graph

The first step of developing a GNN involves defining and designing the graph structure to be used. Formally a graph is defined as \( G = (V, E) \) where \( V \) is a set of nodes, and \( E \) a set of edges. Let \( v_i \in V \) denote a node; then each edge \( e \in E \) is defined as \( e_{i,j} = (v_i, v_j) \), that is pointing from node \( v_i \) to node \( v_j \). In addition to that for a graph with \( n = |V| \) nodes and \( m = |E| \) edges a feature matrix \( X \in R^{n \times d} \) is defined where each node \( i \) carries a feature vector \( x_i \in R^d \). Note that \( d \) denotes the dimension of the feature vector. This encodes the information for the structure represented by the node. Often it is also important to encode specific information for the edge of two nodes. For this we additionally define an edge feature matrix \( X_e \) where \( X_e \in R^{m \times c} \) with each edge \( e \) between nodes \( i \) and \( j \) carrying an edge feature vector \( w_{i,j} \in R^c \) where \( c \) denotes the dimension of the edge feature vector.

Using the above definitions there are two common scenarios, either the data is inherently structured or inherently unstructured. In the first case, a direct translation into the above structure is possible, although additional information may be encoded by the designer. For example in knowledge graphs or molecule simulations such a direct translation would be possible, as the data itself already forms a valid graph, only the feature vectors would have to be engineered in an appropriate way. In the second scenario the data implicitly contains a graph-like structure, as is the case for ATS. Natural language text contains structure but that structure is not directly available as a graph. Simplistically one could consider text as a linear graph, that is each sentence is a node and all sentences following each other are connected with edges. However, this would not expose all the possible information hidden, as text implicitly contains much structural and semantic information, both of which can explicitly be modeled with the help of a graph.

A common scenario for ATS is that the text is encoded into the graph using word nodes and sentence nodes. The above definition does not allow such an option as it assumes an homogeneous graph, that is each node’s feature vector is equal in dimension. However, for ATS and many other tasks heterogeneous graphs are more common. This is due to the fact that heterogeneous graphs allow to include different types of structures within the same graph. However, all of this is not an issue as one can define a feature matrix \( X \) for each type of node that exists within the graph. In the case of ATS with word and sentence type nodes one would define \( X_w \in R^{n \times d_w} \) as the feature matrix for all word nodes and \( X_s \in R^{n \times d_s} \) as the feature matrix for all sentence nodes with \( d_w \) being the dimension of the word node feature vectors, and \( d_s \) representing the dimension of sentence node feature vectors.

For ATS it is also sometimes useful to utilize the design space of directed vs. undirected edges. The choice of directed vs. undirected edges allows to explicitly encode structural information. For instance, one can explicitly encode the order of words or sentences with directed edges.

2.2 Defining the Neural Network

There are three ways in which GNNs differ from more conventional neural networks, the input, the output and the way information is aggregated within the network. The input is the above described graph.

The output and associated loss function are in the case of GNNs bound to the structure of the data i.e. the graph. Generally there are three levels of output possible, namely, on the node level, on the edge level and on the graph level. Both node and edge level output involve predictions or classifications on their respective components of the graph; additionally, in the case of edge level output, one can pose the task of predicting the edge itself. On the graph level one can pose the task of predicting subgraphs or perform graph segmentation. In the context of ATS, the most direct output and loss formulation consist in predicting binary inclusion labels on nodes representing sentences, or phrases, using a cross-entropy loss.

The way information is aggregated is usually done through either spatial convolution or spectral convolution. The basic idea of spatial convolution involves extending the well-known convolution operator to graph structures, whereas spectral convolution is based on graph signal processing. In recent years spatial convolution has become a
popular approach and is the preferred approach for GNNs for ATS due to the flexibility and efficiency it offers. More specifically, in terms of efficiency, spectral convolution involves either computations over the entire graph or eigenvector computations. None of which are necessary for spatial convolution; additionally, spatial convolution does not assume a fixed graph structure, allowing better generalization. In addition to this the locality of the spatial convolution operation also allows it to be performed in batches of nodes, instead of the entire graph. This is especially relevant for large graphs, or, in the context of this paper, large input texts.

The setup described thus far implicitly assumes that the GNN performs extractive summarization. This is in fact the case and for what we will later define as standalone GNNs i.e. GNNs that are not part of a larger system. No purely abstractive approaches have so far been developed, to the best of our knowledge.

### 2.3 Spatial Convolution and Message Passing

One can view spatial convolution as used in GNNs as a generalization of the convolution used in neural networks such as CNNs. As an example, in the case of images, one can imagine 2D convolution as being applied to a regular grid of nodes where each node represents a pixel in the image. The resulting 2D convolution applied to one target node is then the weighted average of node (pixel) values of the neighbours of the target node. Generalizing this idea to a non-regular grids leads to spatial convolution. However, different to images and regular grids, in graphs, the neighbours of each target node are unordered and can vary in number and their feature vector representation. The major challenge with this extension consists therefore in dealing with the unordered and inconsistent neighbourhood sizes inherent to homogeneous and heterogeneous graphs, with an additional challenge being posed by the differing feature vector representations in heterogeneous graphs.

Directly translating the above description of convolution into a mathematical formulation leads to a valid information propagation scheme. However, such a description suffers from scalability issues due to it directly operating over the entire graph. As such modern GNNs use, what is commonly referred to as, message passing. In practice, this means that nodes within the graph exchange messages (perform convolutions) with their neighbours for a number of iterations. Thereby the network is able to diffuse information throughout the graph. Consequently, the more iterations, the further outwards information is propagated throughout the graph. In the terminology of CNNs one would say that the more message passing iterations, the larger the receptive field of the convolution. Formally, one can define message passing (Grattarola and Alippi, 2021) for each time step \( t \) as two equations:

\[
m_{i,j}^{t+1} = \phi(x_i^t, x_j^t, w_{i,j}^t), e_{i,j} \in E
\]

This first equation describes how messages are generated. A differentiable function \( \phi \) generates messages \( m \) for each edge which connects nodes using the node features and edge feature present.

\[
x_i^{t+1} = \psi(x_i^t, \rho(\{m_{i,j}^{t+1}: e_{i,j} \in E\}))
\]

The above equation is the core of the message passing framework and describes how each node feature is updated. The first part consists in the application of a permutation-invariant reduction function \( \rho \). This function aggregates all incoming messages to a node. Then another differen-
tiable function $\psi$ combines the reduced messages received with the previous state. Using these two equations one can utilize message passing for learnable layers.

The convolution layer for a GNN is then defined with a learnable weight $W$ such that the message per edge is $m_{i,j}^{t+1} = x_i^t$, and the aggregation is the normalized sum of messages, i.e. $x_i^{t+1} = \sigma(b^t + \sum_{m_{i,j} \in M(i)} \frac{1}{c_{i,j}} m_{i,j}^{t+1} W^t)$ where $M(i)$ represents the set of messages received by node $i$, $\sigma$ is the activation function, $b$ is the bias, and $c_{i,j}$ is an appropriate scaling factor, e.g., the square root of the node degree. Note how it is important for the reduction function, in this case a sum function, to be permutation-invariant as otherwise GNNs could not handle the unordered nature of graphs.

The above presented convolution layer does not allow the model to filter unimportant neighbours. Inspired by the attention mechanism popularized by transformer networks (Vaswani et al., 2017), graph attention networks (GAT) (Veličković et al., 2018) assign attention scores to each neighbour. A schematic depiction of the two variants of spatial convolution can be seen in Figure 1 with GAT depicted on the lower part of the figure. The introduction of attention scores to the spatial convolution allows the model to explicitly assign importance to certain nodes and their messages. Just as in transformers GAT is formulated with multi-head attention. The modification to the previously presented convolution layer follows closely the common attention formulation. Formally, $x_i^{t+1} = \|_K \sigma(b^t + \sum_{m_{i,j} \in M(i)} \alpha_{i,j} m_{i,j}^{t+1} W^t)$ where $\alpha_{i,j}$ is the attention score between node $i$ and node $j$ and $K$ denotes the number of concatenated heads. The attention scores are computed with $r_{i,j} = \text{LeakyReLU}(a^T [W x_i^t \| W x_j^t])$. This score is then normalized to obtain the attention score per edge $\alpha_{i,j} = \text{softmax}(r_{i,j})$. We want to highlight here a recent development which Brody et al. (2021) simply denote as GATv2. Their main improvement aims at the fact that in the above calculation both learnable parameters $a$ and $W$ effectively fold into a single linear layer, thus the expressive power of the layer is less than what it could be. The fix introduced by GATv2 pulls the two parameters apart, thus achieving more expressive power while not increasing computational complexity. Taking the above description the attention score for GATv2 is modified as follows $r_{i,j} = a^T \text{LeakyReLU}([W x_i^t \| W x_j^t])$. In both synthetic and real datasets this modification shows superior performance, which is supported by a theoretical analysis of the authors.

There are numerous modifications and extensions to the basic convolution presented here. However, for ATS models, GAT layers are dominating as the workhorse for most models. The reasoning for their dominance can be explained by the similar success that attention transformers have had in conventional neural networks for AST. We expect that GATv2 will continue this trend as it is an attractive and simple improvement for the currently dominating GAT. Although the authors of GATv2 note that it is not yet entirely clear which tasks would benefit the most from the usage of GATv2 over GAT, which will require more research and models to use GATv2.

In ATS the graphs used are in nearly all cases not homogeneous. However, the equations presented here do not work for heterogeneous graphs. The solution for this problem involves defining one convolution layer for each node type combination occurring within the graph. In the case of the already discussed sentence and word node graph there would be four possible combinations of types, and four convolution layers which would have to be defined if the graph were fully connected.

Convolution is a central aspect of GNNs, but pooling also presents an important and common operation, especially whenever GNNs are used jointly with other models. Pooling in GNNs is achieved by generating a global representation of the graph, or a subset of the graph, by pooling together features of nodes. This is usually done with some function $f$ where $f$ is commonly the mean, max or sum.

We want to explicitly point out to the reader that the construction of GNNs does not require special datasets. All GNN models for ATS use the common benchmark single-document and multi-document summarization datasets such as DUC 2004 or CNN/Dailymail. The only requirement for any ATS, or textual dataset, is for the designer to find an appropriate way of encoding sentences, words, subwords etc. into feature vectors, and finding a sensible way of connecting them.

3 Graph Neural Network Models For Automatic Text Summarization

The current state of research clearly shows that the usage of GNNs for ATS follows one of two patterns,
either the GNN is used directly for the ATS task or it is used to support a larger system. These trends justify the taxonomy as seen in Figure 2. More precisely, we classify standalone ATS GNNs as GNN models which are directly responsible for generating a summary. Embedded GNNs on the other hand are used to support a larger system, and are not directly or solely responsible for producing the summary. In addition to this difference, we further differentiate between embedded GNNs used in an encoder-decoder setting and other embedded GNNs.

Note that we do not specifically differentiate between single and multi-document GNN summarization models as some models are by design capable of handling both tasks. Additionally, we do not make a distinction between abstractive and extractive approaches, since embedded GNNs can be part of either approach, while abstractive approaches have not yet been developed for standalone GNNs, to the best of our knowledge.

3.1 Standalone GNNs

We will start our discussion of standalone GNN models with HeterSumGraph (HSG), a model proposed by Wang et al. (2020). We will do so due to the fact that this model illustrates concepts and ideas seen throughout GNNs models used for ATS. An illustration of the general concepts presented here can be seen in Figure 3.

The HSG model encodes each text into a graph with three node types, sentence nodes, word nodes and document nodes. The connection between these nodes is decided by inclusion i.e. if the word represented by a word node occurs in a sentence then their respective nodes are connected by an edge. The same principle applies to document nodes which are connected depending on whether a word, represented by a word node, occurs within the document. This is a flexible structure, as it can be used in a single-document but also multi-document setting.

The feature vectors for all nodes are obtained by encoders and the edge weights are obtained by computing the TF-IDF score for each word. The neural network consists of a modified GAT layer. The GAT is modified to consider the TF-IDF value of the connecting edge. Additionally a position-wise feed-forward (FFN) layer consisting of two linear transformations is applied to the hidden state after the convolution. In total three convolution layers are used, word-sentence, sentence-word and word-document. The model is then trained on a node-based binary classification task that is predicting whether a sentence node is to be included for the summary or not.

The classification itself is done by a single linear layer. The model then does not directly use the predicted nodes to produce the summary. Instead trigram blocking (Paulus et al., 2018) is used during sentence selection in order to ensure sparsity of the generated summary.

The results for this model are quite impressive as it outperforms non-BERT based models on both single-document and multi-document summarization for the CNN/DailyMail dataset. One should especially note the flexibility and ability to use this model for two tasks.

An older model by Muratore et al. (2010) can be considered a precursor to this architecture. A simple extension to the HSG model is proposed by Ya et al. (2021). In their extension they modify the model for query constraints for the summary. This is achieved by adding a query node to the graph structure. Additionally, they introduce a mu-
tual information maximization mechanism during training.

A model which further follows this structure is the one by Linmei et al. (2019). The authors there extend the attention mechanism by adding another layer of attention, allowing it to include information about the type of the node during convolution. The GNN model by Jing et al. (2021) encodes even more information into the graph by considering the relation between sentences on a number of different levels. In particular, they encode the semantic and syntactical relationship between sentences within the graph.

This idea of encoding additional information into the graph is also followed by Antognini and Faltings (2019). They introduce an additional universal feature vector which is added to each sentence node embedding. This universal feature vector is learned from a large unrelated and general corpus. This model is also unique in that it focuses on the summarization of very small texts, on average less than 100 words.

Taking this basic structure and idea even further is the model called HAHSum by Jia et al. (2020a). The construction of the input graph for HAHSum is more involved as it aims to significantly reduce semantic sparsity by utilizing named entities. The model uses three types of nodes, named entity nodes, word nodes and sentence nodes, with the named entity nodes being anonymized tokens. The graph is then built as follows, word nodes are connected with a directed edge to a sentence node if they occur within the sentence. Two named entities are connected with an undirected edge if they represent the same entity and two sentence nodes are connected with an undirected edge if they share a trigram. Additionally, sequentially occurring words and entities are connected with a directed edge. This setup shows how one can encode a substantial amount of implicit information in an explicit manner.

HAHSum uses a GAT for each of the five node type combinations found within the graph. Just as in HSG, a FFN is applied after the multi-head attention and again as in the previous model a linear layer is used to perform the binary classification of the sentence nodes.

The results for HAHSum show that GNNs can perform very well. The authors of the paper tested the model on the CNN/Daily Mail, Newsroom and NYT dataset. The model outperforms very powerful models such as MATCHSUM (Zhong et al., 2020) and is even able to compete in some metrics with leading abstractive models such as PEGASUS (Zhang et al., 2020a). The results of an Amazon Mechanical Turk experiment corroborate these results and show that for human readers HAHSum produce summaries with superior fluency and conciseness.

Another recent GNN model which has achieved great performance in the task of multi-document summarization is the SgSum model by Chen et al. (2021). Different to the approaches outlined above, the SgSum model uses graph pooling to extract sub-graphs from encoded documents. That is, it first transforms the documents into a large graph, then generates a number of sub-graphs via pooling and convolution. These sub-graphs are then ranked and thereby selected for a summary. This is quite an innovative approach as it casts the problem of multi-document summarization as a simple sub-graph selection problem. Additionally, it outputs an integral summary, that is the entire summary is output by the model in the form of the sub-graph of sentences.

3.2 Embedded GNNs

The first embedded GNN model we will present is the GRU-GCN model by Yasunaga et al. (2017). Despite being an older model it provides an illustrative introduction as to how one can effectively incorporate GNNs into established deep learning methods. As this model utilizes GNNs within a sequence-sequence architecture it falls into the encoder-decoder category of embedded GNNs. The model works exclusively with multi-document summarization. We want to note that due to the age of this model the GNN uses spectral-based convolution instead of spatial-based convolution.

As the model is an encoding-decoding embedded GNN it features three parts: the encoding part, the GNN and a decoding part. The encoding and decod-
Table 1: Combined results as reported in their respective papers. We only report the best performing model type and only ROUGE-1 (R-1), ROUGE-2 (R-2) and ROUGE-L (R-L) scores. We show results on the CNN/Dailymail dataset for single-document summarization and results on the DUC-2004 dataset for multi-document summarization. NA denotes that the authors have not reported this score. Best results marked with boldface.

| Model Name | Type | Description | R-1 | R-2 | R-L |
|------------|------|-------------|-----|-----|-----|
| BERTSUMEXT (Liu and Lapata, 2019) | Extractive BERT baseline | BERT-based | 42.85 | 20.34 | 39.90 |
| Topic-Graphsum (Cui et al., 2020) | Extractive (Embedded) | NTM + GNN | 44.02 | 20.81 | 40.55 |
| DSGSum (Bi et al., 2021) | Abstractive (Embedded) | Seq-Seq with GNN | 41.96 | 19.29 | 38.98 |
| Syntactic-Graph (Xu et al., 2020a) | Abstractive (Embedded) | Syntactic graph | 41.79 | 19.06 | 38.56 |
| HABSUM (Jia et al., 2020b) | Extractive (Standalone) | Entity-relations + words and sentences | 44.68 | 21.30 | 40.75 |
| Multi-Gras (Jing et al., 2021) | Extractive (Standalone) | Sentence information encoding | 43.16 | 20.14 | 39.49 |
| HSG (Wang et al., 2020) | Extractive (Standalone) | Multi-layer encoding of documents | 42.95 | 19.76 | 39.23 |
| DISCOBERT (Xu et al., 2020b) | Extractive (Embedded) | BERT+GNN | 43.77 | 20.85 | 40.67 |

| GNN DUC-2004 Performance Overview |
|-----------------------------------|---|---|---|
| GRU+GCN (Yasunaga et al., 2017)  | Extractive (Embedded) | GRU+GNN | 38.23 | 9.48 | NA |
| SGGSum (Chen et al., 2021)       | Extractive (Standalone) | Subgraph extraction | 39.41 | 10.42 | 35.41 |

The authors of (Wu et al., 2021b) produce a graph based on the semantics of each sentence. The encoding produced by the GNN and a textual encoding of each sentence are then passed to a decoder. Similarly, the model by Xu et al. (2020a) uses a GNN as an encoding component. Their model utilizes the dependency tree of the input as a graph input to the GNN. It also uses a modified attention mechanism which is used to decode the attention of the GAT directly into the decoder part of the architecture. Another model in this category is the model by Liang et al. (2021). Following this idea even further is the model by Li et al. (2020). However, in contrast to DSGSum or others, their model uses a GNN directly within the transformer based encoder and decoder blocks, and not as an outside component providing additional encodings.

Another area where GNNs have found some usage is in the abstractive summarization of di-
alogues, which although a niche area, is still part of the ATS task. The model by Zhao et al. (2020) uses a GNN to encode the structure of the conversation into their sequence-to-sequence architecture. This is also done by Feng et al. (2020). Also for dialogue summarization Feng et al. (2021) introduce a special type of graph encoding to a sequence-to-sequence architecture. They utilize a GNN as an encoder and their input graph links information about the speaker, the spoken sentences and other information together.

At this point we also want to shortly mention a few models which do not perform classical ATS but do perform a specialized form of summarization with the help of a GNN. The models by Liu et al. (2020a) and LeClair et al. (2020) both perform code summarization, that is they generate a natural language description/summarization of code written in a programming language. They both utilize a GNN, and also both leverage the abstract syntax tree of the program given. Another interesting model is the model by Wu et al. (2019). Their model performs multi-video summarization with the help of a GNN.

4 Conclusions and Outlook for Future

We have surveyed in this paper the main developments in the area of GNNs applied to the automatic text summarization task, first describing how GNNs work and then discussing the prominent models used for ATS. We have also provided simple categories of the models. Finally, in Table 1 we have combined results from the models mentioned in our survey. Note that this is just a simple enumeration of the best ROUGE scores reported, which does not fully capture other important aspects of summaries such as fluency, conciseness, relevancy and in the case of abstractive summarization, factual accuracy.

We have already highlighted some general advantages of GNNs for ATS. Now considering the models presented and the results shown in Table 1 we want to give some pointers towards the future of GNNs for ATS.

- **Outside the Box.** Models such as SGSum and HAHSUM show how one can achieve great performance by rethinking parts of the ATS task. GNNs provide with their direct acceptance of graphs a lot of freedom with regards to the design of the input as well as the output. The idea of reconsidering multi-document ATS as a subgraph ranking task is a worthwhile approach to consider further.

- **Encoding and Enhancing.** GNNs are able to efficiently and effectively produce encodings of graph structures, which are prevalent throughout texts. Models like DISCOBERT show how one can enhance the performance of traditional models by incorporating GNNs and leveraging their encoding ability. We believe many models can be improved by incorporating GNNs in ways presented in the survey.

- **Text Length.** GNNs are capable of scaling to very large graph sizes. Summarizing large texts is an important task, but one which has been neglected in the literature. GNNs are in our estimation in a position to perform very well on large text summarization.

- **Explainability.** No GNN ATS model so far has leveraged the explainability aspect of GNNs. We believe that this is a very unique and unexplored research avenue that could reveal valuable insights into ATS and GNNs used for ATS.

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