SELF-LABELING OF FULLY MEDIATING REPRESENTATIONS BY GRAPH ALIGNMENT

Martijn Oldenhof, Adam Arany, Yves Moreau & Jaak Simm
ESAT - STADIUS
KU Leuven
Leuven, 3001, Belgium
{martijn.oldenhof,adam.arany,yves.moreau,jaak.simm}@esat.kuleuven.be

ABSTRACT

To be able to predict a molecular graph structure \( W \) given a 2D image of a chemical compound \( U \) is a challenging problem in machine learning. We are interested to learn \( f : U \rightarrow W \) where we have a fully mediating representation \( V \) such that \( f \) factors into \( U \rightarrow V \rightarrow W \). However, observing \( V \) requires detailed and expensive labels. We propose graph aligning approach that generates rich or detailed labels given normal labels \( W \). In this paper we investigate the scenario of domain adaptation from the source domain where we have access to the expensive labels \( V \) to the target domain where only normal labels \( W \) are available. Focusing on the problem of predicting chemical compound graphs from 2D images the fully mediating layer is represented using the planar embedding of the chemical graph structure we are predicting. The use of a fully mediating layer implies some assumptions on the mechanism of the underlying process. However if the assumptions are correct it should allow the machine learning model to be more interpretable, generalize better and be more data efficient at training time. The empirical results show that, using only 4000 data points, we obtain up to 4x improvement of performance after domain adaptation to target domain compared to pretrained model only on the source domain. After domain adaptation, the model is even able to detect atom types that were never seen in the original source domain. Finally, on the Maybridge data set the proposed self-labeling approach reached higher performance than the current state of the art.

1 INTRODUCTION

Chemical compounds are often represented by a graph representation of their chemical structure. These graph representations are actually a simplification of the chemical compound as it loses some information about the electronic structure of the molecule. However, in the field of drug discovery this graph representation is often used as valuable input for machine learning pipelines. Examples of formats describing the graph representation of a chemical compounds are SMILES \(^{30}\) and MOLfile \(^3\). However, especially in patents but also in scientific literature the chemical compound is only described using an image format. Automatically recognizing the chemical structures on these images is valuable for machine learning approaches to be able to process these sources of chemical compounds.

Learning to recognize a graph structure from 2D images of chemical compounds seems like a fairly simple task for humans. However, for machine learning models it seems that generalization to new domains of images (e.g. different line width, font face) \(^{21}\) is not happening naturally. When we humans see an image with a graph structure that we do not recognize completely, we start reasoning and analyzing the part of the graph we are not sure about. We humans automatically align the graph part we recognized on the image with the complete graph including the unrecognized part of the graph. One way to finish our graph prediction is to guess the unknown nodes or edges after which

\( ^{1}\) Code available: https://github.com/biolearning-stadius/chemgrapher-self-rich-labeling .

arXiv:2103.14133v1 [cs.LG] 25 Mar 2021
we check for correctness. If the graph prediction was correct we know that this guess was most probably correct and we could try to apply this new knowledge to other images.

To be able to do this reasoning on for example images using graph alignment in machine learning we need a detailed (on pixel level) representation. Therefore we assume a fully mediated model \( f : U \to W \) where we are interested to learn \( f \) factors into \( U \to V \to W \), which is visualized in Figure 1. Thus, in order to predict \( W \) from \( U \) we first need to pass the fully mediating layer, no side paths are allowed. When a fully mediating representation is used some assumptions \([23; 26; 25]\) are made about the mechanism of the underlying process. This mechanistic prior restricts the space of possible models to all the models that follow the mechanistic assumption. We hypothesize that the use of this richer representation (fully mediating representation) enables for a better generalization. Additionally, as an interesting side effect, we observe that the mechanistic assumption allows for a better interpretability of the underlying model.

In the case of optical graph recognition of chemical compounds from 2D images, the fully mediating layer is represented using the planar embedding of the chemical graph structure we are predicting. In order to learn the planar embedding of a chemical graph structure, we start from a model described in Oldenhof et al. \([21]\) which has two steps: an image segmentation and an image classification step. To train this model, pixel-wise labels are needed for every image describing precise locations of nodes and edges in the graph (planar embedding) which we will call rich or detailed labels in our setup. However, these rich labels are not always available and implies a manual process where intermediate organic chemistry knowledge is required. In the more common cases, data sets only contain 2D images of chemical compounds (\( U \) in Figure 1) and on the other side the final output in SMILES\([36]\) or MOLfile\([5]\) format (\( W \) in Figure 1). These formats describe the graph structure of the chemical compound but not the particular planar embedding of this graph structure (\( V \) in Figure 1) in the context of the image. To solve this problem, we propose a graph aligning approach that generates rich labels \( V \) given normal labels \( W \). This method would enable learning of the fully mediating representations given only normal labels \( W \).

In section 4 we empirically evaluate our domain adaption method. We observe that compared to the non-adapted model we drastically increase accuracy even on atoms and bond that were not present in source domain.

**Key contributions:** (1) we propose a novel rich labeling framework by introducing the use of fully mediating representations, (2) in the case of graph recognition we show that the rich labeling can be performed by graph alignment, (3) we show it enables data efficient domain adaption and (4) reaches state-of-the-art performance on Maybridge compound data set.

## 2 RELATED WORK

**Structural scene representation and visual reasoning.** Our work has similarities with research done on structural scene representation and visual reasoning \([11; 41; 19]\). The disentanglement of the reasoning and the representation described in Yi et al. \([41]\) enables the model to solve complex reasoning tasks. In our work the complex reasoning task would be graph alignment which is disentangled from the optical graph recognition.

**Slot Attention.** Our method is related with a method called slot attention \([17]\) where the Hungarian algorithm \([16]\) is incorporated in a model for object detection. This Hungarian algorithm is limited to only sets while in our case we need to map more complicated structures composed of different atoms connected with different bond for which we need graph alignment in order to adapt iteratively a model to a new target domain.

**Image to Graph methods.** In the field of computational chemistry there are several tools available \([22; 34; 20; 7; 21; 18; 33]\) to convert an 2D image of a chemical compound to a SMILES \([36]\) format or similar which in fact represents a graph structure of a chemical compound. Also for road extraction from satellite images there are several methods available \([3; 8; 12]\).

**Graph Matching.** In computer vision graph alignment is usually known as graph matching. It can be useful to (1) locate objects from features \([10]\), (2) to transfer knowledge \([42]\) and (3) to find matches in database \([13]\). Also for comparing social networks graph matching can be very important.
Figure 1: We are interested to learn \( f : U \to W \) having a fully mediating representation \( V \) such that \( f \) factors into \( U \to V \to W \). In the case of optical graph recognition of chemical compounds from 2D images, the fully mediating layer is represented using the planar embedding of the chemical graph structure we are predicting.

to allow to uncover identities of communities \[14\]. In chemistry, comparing graphs can be helpful to identify identical chemicals, substructures or maximum common part of chemicals. In the work of Willett et al. \[37\] an overview is presented about the use of similarity searches in chemical databases.

**Domain Adaptation** In the work of Kouw & Loog \[15\] a comprehensive overview is given for domain adaptation methods when labels for the target domain are not available. Our method has some similarities with semi-supervised iterative self-labeling \[4, 27\] approaches where predictions on a data set of a new domain of a pre-trained model are used as pseudo-labels and used to retrain the model again iteratively until convergence. In the work of Das & Lee \[6\] even a graph matching loss is first used to learn a domain invariant representation for source and target domain after which the use of pseudo-labels show a significant improvement of performance. In our work the graph matching is used for a different purpose as opposed to the the work of Das & Lee \[6\]. Graph matching is used in our work to generate rich labels given the 'normal' labels we have from the target domain. This is where our method also differs from other semi-supervised methods for domain adaptation when no target label information at all is assumed and no distinction is made between rich and 'normal' labels.

**Weak Supervision** In our setup we use the term 'rich' or 'detailed' labels to differentiate from the normal labels. We would like to contrast these ‘rich’ labels with the term ‘strong labels’ used in the setting of weak supervision. For example, in the machine learning task of image segmentation pixel-wise labels are needed which are expensive and often not readily available. Therefore, weak supervision methods have been developed to address this issue. Weak supervision can be used to help image segmentation by only using image labels (no pixel-wise labels) \[35, 38\]. A more general framework was presented in Xu et al. \[39\] to be able to learn semantic segmentation from a variety of types of weak labels (e.g., image tags, bounding boxes and partial labels). Another approach is to augment the strong labeled data set using weakly labeled data \[40\]. However, the main difference with all of the methods mentioned above is that our method does not work on weak labels because our end goal is different. The main goal of our machine learning approach is to help to predict ‘normal’ labels by using rich labels.

**Front Door Criterion** Our framework exploits fully mediating variables. A variable is called a mediator when it meets several conditions regarding the relationship with other variables as described in Baron & Kenny \[2\]. Another perspective of the mediating relationship is given by Pearl \[23\]; Pearl et al. \[20\]; Pearl \[25\] who introduce the front door criterion where the mediator actually enables to estimate unbiased causal effects. A more formal interpretation of these causal effects is presented in Pearl \[24\]. In order to use a mediating model the mediator needs to be identified or assumed
Figure 2: Two examples of chemical compounds graphs with their graph edit distance. The nodes of the graphs are first aligned before computing the graph edit distance. The node alignments are marked with the gray dashed arrows. The differences after graph alignment are highlighted and the substitutions are marked with the red dashed arrows.

first, which is not always straightforward. In our setup (see Figure 1), the assumption means that the relation between input \( u \in U \) and planar embedding \( v \in V \) is a map and as well as the relation between \( v \) and the final graph \( w \in W \). Furthermore, we assume no side paths from \( u \) to \( w \).

3 SELF-LABELING OF FULLY MEDIATING REPRESENTATIONS

Our goal is to learn \( f : U \rightarrow W \) assuming a fully mediating representation \( V \) such that \( f \) factors into \( U \rightarrow V \rightarrow W \). In order to learn the first part of \( f (U \rightarrow V) \) we need labels for \( V \) which are expensive in the case of optical graph recognition of chemical compounds from 2D images where \( V \) is represented as the planar embedding of the chemical graph structure. Our method tries to address this issue by iteratively updating the model using self-labeled labels for \( V \) by graph aligning the graph predictions using the model from previous iteration with the given true graphs (labels \( W \)).

3.1 GRAPH ALIGNMENT

A possible and often used closeness score to compare graphs is the **graph edit distance** [29]: given 2 graphs, not necessarily of equal size and a set of operations, that are \( \mathcal{O} = \{ \text{vertex/edge/label insertion/deletion/substitution} \} \), and a cost function \( c : \mathcal{O} \rightarrow \mathbb{R} \), so we find the cheapest sequence of operations that convert \( G_1 \) into \( G_2 \), which translates to an optimization problem:

\[
\min_{\{e_i\}_{i=1}^k \in \mathcal{O}^k : G_2 = (e_k \circ \ldots \circ e_1) \times G_1} \sum_{i=1}^k c(e_k),
\]

Although there are some efficient algorithms available \([30, 31, 32]\) in order to compute the graph edit distance, it remains a computational hard problem.

Closely related with the concept of graph edit distance we introduce for our method the map \( E(v) \) which gives the allowed operations on a given graph \( v \) given a specific constraint. This constraint is a parameter which can be tuned for a specific data set or problem domain. Examples of such constraints are maximum 2 node substitutions or maximum 1 edge substitution as shown in Figure 2.

3.2 METHOD

Let us now say we have a trained neural network model for \( f : U \rightarrow V \), a projection \( \phi : V \rightarrow W \), a pair \((u, w)\) of input \( u \in U \) and normal label \( w \in W \) and we would like to infer rich label \( v \in V \) from the given datapoint \((u, w)\). We also assume the map \( E(v) \) which gives all allowed graph edits for the graph \( v \). Let \( \hat{v} = \hat{f}(u) \) be the predicted rich label from the model, then we define a term correcting edit as
**Definition 3.1** Edit $e$ is a **correcting edit** if when $e$ is applied to the prediction $\hat{v}$ and then projected to the $W$ space the resulting graph is the true graph $w$ (up to isomorphism), i.e.,

$$\phi(e \times \hat{v}) \cong w,$$

where $\times$ is the application of edit to the planar embedded graph $\hat{v}$.

Notice that for a given $\hat{v}$ and $w$ there can be multiple edits that are **correcting edits** which create a dilemma of choosing the best correcting edit. Therefore, we make the following assumption:

**Assumption 3.1** The probability that a correcting edit $e$ results in the true underlying rich label $v$ is monotonely decreasing with respect to the size of edit $e$ (i.e., $|e|$).

In other words, if we take two correcting edits $e_1, e_2$ then we assume the following:

$$|e_1| < |e_2| \implies P(e_1 \times \hat{v} = v) > P(e_2 \times \hat{v} = v)$$

The assumption is based on the fact the probability of any individual mistake in a graph by the model is low. This is because if the probability of a mistake would be high the model would not be able to produce a graph with a total of 1-2 edit distance. Thus, the graphs with few edits have low mistake probability and for them the Assumption 3.1 is valid.

Then we use the following optimisation problem to find the best correcting edit $e$ to convert $\hat{v}$ to rich label $v$ for input $u$:

$$\mathcal{E}^* = \arg \min_{e \in E(\hat{v})} |e| \text{ such that } \phi(e \times \hat{v}) \cong w,$$

where $\arg \min$ returns the set of minimal solutions or the empty set if no solutions exist.

There are three possible outcomes of last mentioned optimization problem: (1) no solution is found, (2) a single $e$ is found or (3) multiple equal size $e$ are found. In the optimal case (2) a single $e$ is found so we can label a new $v$ for our given datapoint $(u, v)$. In the case of (1) when no solution is found, no new $v$ is labeled. In the last case (3) when multiple equal size solutions are found there are four options we could do. First (3.1), we could discard the solutions and not label $u$. Second (3.2), we could take $e$ that results in the highest likelihood for $e \times \hat{v}$ based on the model $f$. Third (3.3), a solution $e$ is picked uniformly randomly in order to generate the rich label label $v$. Fourth (3.4), pick $e$ randomly according to the likelihood of $e \times \hat{v}$ in the model $f$.

This process is repeated for every datapoint $(u, w)$ we have available from the target domain. Thus, several new labels $v$ are found for different datapoints. Once all datapoints are processed these new rich labeled datapoints are added to the training data set after upsampling and our model can be retrained. After this, a new iteration begins and all available datapoints $(u, v)$ are again processed to...
Table 1: Summary of datasets from the 2 different target domains

| Dataset  | Orig. Size | Tot. #samples | Cand. rich labeling #samples | Test #samples |
|----------|------------|---------------|-----------------------------|--------------|
| Indigo   | 50,000     | 5,000         | 4,000                       | 1,000        |
| Maybridge| 5,740      | 5,000         | 4,000                       | 1,000        |

find even more new rich labels $v$ and we can retrain the model again. This iterative process can be repeated until convergence (see Algorithm 1).

Algorithm 1: Iterative algorithm for Self-Labeling of Fully Mediating Representations

Data:
Target domain data $L = \{(u^i_t, w^i_t)\}_{i=1}^n$
Source domain data $S = \{(u^j_s, v^j_s)\}_{j=1}^m$ (rich labels)

Result: $f : U \rightarrow V$

repeat
  // Inferring rich labels for target data
  $T = []$;
  for $(u, w)$ in $L$ do
    $\hat{v} \leftarrow f(u)$;
    $E^* \leftarrow \arg\min_{e} |e|$ such that $\phi(e \times \hat{v}) \cong w$;
    if $E^*$ is a not empty then
      $e \leftarrow \text{choose}(E^*)$;
      $v \leftarrow e \times \hat{v}$;
      appendRichLabels($T$, $(u, v)$);
    end
  end
  $T \leftarrow \text{UpSample}(T)$;
  $f \leftarrow \text{RetrainModel}(S, T)$;
until Converged($f$);

4 Experiments

For the experiments we focus on the problem of predicting chemical compound graphs from 2D images where the fully mediating layer is represented using the planar embedding of the chemical graph structure we are predicting. In order to measure empirically the performance of our method of self-labeling fully mediating representations we perform three steps. (1) We pre-train (training details in Appendix A.2) a ChemGrapher [21] model (summarized in Appendix A.1) wherefore, corresponding to the pipeline described in the work of Oldenhof et al. [21], we sample around 130K chemical compounds from ChEMBL [9] in SMILES format and artificially generate, using an RDKit fork [1], a rich labeled dataset with 2D images of chemical compounds. (2) Secondly, we test the baseline performance of this pre-trained model on two different test sets from two different target domains than the source domain of the pre-trained model. (3) Thirdly, we apply our domain adaptation method and measure performance again on the two target domains.

For the first target domain we take a data set from the work from Staker et al. [33], which we will call Indigo data set. For the second target domain we take the data set which was published by the developers of MolRec [18] which we will call the Maybridge data set. Both data sets provide 2D images from a chemical compound together with corresponding identifier of a the chemical compound like SMILES [36] or MOLfile [5]. These identifiers describe the graph structure of the chemical compound however they do not provide the planar embedding of the graph (e.g. no information about the pixel coordinates of every node or edge in the image). Visually we can also observe that the Maybridge dataset contains images where the style is closer related to the training images style used for the pre-trained model compared with the images in Indigo dataset where the style of images is quite different. Therefore we expect a significant worse starting performance of the pre-trained model on the Indigo dataset compared with the Maybridge dataset.
(a) Results on Indigo data set

(b) Results on Maybridge data set

Figure 3: Comparison performance of methods on Indigo and Maybridge data set. Self-labeling by correcting graph alignment is clearly better performing than when exact graph alignment is used. Sometimes upsampling of rare atoms to a specific threshold (note postfix \textit{rare}) before retraining of model can boost performance. Performance on target domain at iteration 0 is the performance of pre-trained (on source domain) ChemGrapher before domain adaptation.

From both data sets we randomly sample 5,000 datapoints which are split in 4,000 datapoints used for our method and 1,000 datapoints to measure performance on (summarized in Table 1). When processing the 4,000 datapoints our method will be able to generate rich labels for the datapoints where the graph prediction could be graph aligned with the true graph. As the number of rich labeled datapoints this way is maximum 4,000 we will upsample them (x number of copies) before adding them to the training data set. In our experiments we differentiate between two strategies of upsampling. One way is to upsample all the rich labeled data points equally from the target domain to a fixed number, for example 20,000. Another way is to take into account, while upsampling, the number of atom types that are rich labeled and make sure that the rare atom types are upsampled to a specific threshold.

One important tuning parameter in our method is the number of allowed operations. For our experiments we will try two different values for this parameter. Firstly, we set this parameter to zero meaning we do not allow any operation for graph alignment. We will call this exact graph alignment. Secondly, we allow a maximum of 2 node substitutions or a maximum of 1 edge substitution for graph alignment, which we will call correcting graph alignment.

In total we will measure the performance of 4 variations of our method (varying allowed operations and upsampling strategy) on both data sets. The performance we will measure is the accuracy of $U \rightarrow W$ as we only have access to the normal labels of target domain. However, we assume that if the final graph prediction is correct ($W$) it is highly likely that also the planar embedding ($V$) is correct. As our method is an iterative method we will report results for every iteration starting with the initial performance before applying our method. The results of these experiments are summarized in Figure 3. We observe that all variations of our method are able to improve performance on target domain compared with initial pre-trained model on source domain. On the Indigo data set the best variation is even able to obtain 4x improvement. The best variation of our method on the Indigo data set was using correcting graph alignment without upsampling of rare atom types while on the Maybridge data set the best variation was also using correcting graph alignment but with upsampling of rare atom types. Some of the underperforming variations of our method were stopped early in order to save computational resources.

We choose the best variation of our method for every data set and analyze the performance on different atom and bond types per iteration. We measure for every atom or bond type the percentage of graphs predicted correctly from the total number of graphs containing that specific atom or bond.
Figure 4: We take the best performing methods and analyze their performance on different atom and bond types per iteration. We observe that for some atom types the method is able to increase performance even though initial performance was 0%. This is the case in for example R-groups in Indigo data set or superatom NO$_2$ in Maybridge data set.

Figure 5: Comparison initial segmentation with final segmentation after applying self-labeling of fully mediating representations for Indigo data set. We observe that the initial model is making mistakes on the R-group atom type and carbon represented with a ‘C’. In the final model we see that now predictions are all correct.

Figure 6: Comparison initial segmentation with final segmentation after applying self-labeling of fully mediating representations for Maybridge data set. The initial model predicts the superatom NO$_2$ as two separate atoms O and N which is chemically not correct. The final model makes the correct prediction.
Table 2: Comparison performance on Maybridge data set. We observe that our approach enables to reach higher performance than the current state of the art. Most of the tools available for chemical graph recognition are rule based approaches for which a training dataset is not relevant.
5 Conclusion

Machine learning models often are faced with the problem to not generalize well to a new domain. This is also the case for chemical graph recognition from images. We have shown that fully mediating layers can be exploited in machine learning models to adapt in data efficient way to new domains, without the need of rich expensive labels as they can be generated using our method. In the case of chemical graph recognition we empirically show that our method is able to adapt to a new domain of chemical compounds, with previously unseen atom or bond types. Our rich-labeling method required only 4,000 normal labeled points in the target domain to go from 10% accuracy to 39%, i.e., almost 4x improvement in the difficult Indigo data set. Furthermore, on Maybridge data set, again using only 4,000 images, we reached high accuracy obtaining better performance than the current state of the art.

Effective tools of chemical structure recognition from images enable access to the knowledge in chemical literature which is currently only available through expensive chemistry databases. We believe it as an important step towards open pharmaceutical science.

It would be interesting to apply this method to other contexts where the output of a machine learning model could be represented with a graph structure. For example, the case of structural scene representation, where a scene could be represented using a graph where every vertex could represent an object and every edge would represent the relations between the objects (e.g. side-by-side, on-top-of, under). This structural scene could be in form of 2D images or it could be even generalized to 3D models, where point clouds are available and one is interested to transform them into 3D graphs of connected parts.

Acknowledgments

MO, AA, YM, and JS are funded by (1) Research Council KU Leuven: C14/18/092 SymBioSys3; CELSA-HIDUCTION, (2) Innovative Medicines Initiative: MELLODDY, (3) Flemish Government (ELIXIR Belgium, IWT: PhD grants, FWO 06260) and (4) Impulsfonds AI: VR 2019 2203 DOC.0318/1QUATER Kenniscentrum Data en Maatschappij. Computational resources and services used in this work were provided by the VSC (Flemish Supercomputer Center), funded by the Research Foundation - Flanders (FWO) and the Flemish Government – department EWI. We also gratefully acknowledge the support of NVIDIA Corporation with the donation of the Titan Xp GPU used for this research.

References

[1] Fork of the official sources for the rdkit library. URL https://github.com/biolearning-stadius/rdkit accessed on 12.11.2020.
[2] Reuben M Baron and David A Kenny. The moderator–mediator variable distinction in social psychological research: Conceptual, strategic, and statistical considerations. Journal of personality and social psychology, 51(6):1173, 1986.
[3] Davide Belli and Thomas Kipf. Image-conditioned graph generation for road network extraction. arXiv preprint arXiv:1910.14388, 2019.
[4] Lorenzo Bruzzone and Mattia Marconcini. Domain adaptation problems: A dasvm classification technique and a circular validation strategy. IEEE transactions on pattern analysis and machine intelligence, 32(5):770–787, 2009.
[5] Arthur Dalby, James G. Nourse, W. Douglas Hounshell, Ann K. J. Gushurst, David L. Grier, Burton A. Leland, and John Laufer. Description of several chemical structure file formats used by computer programs developed at molecular design limited. J. Chem. Inf. Comput. Sci., 32(3):244–255, 1992. URL https://pubs.acs.org/doi/abs/10.1021/ci00007a012
[6] Debasmit Das and CS George Lee. Graph matching and pseudo-label guided deep unsupervised domain adaptation. In International conference on artificial neural networks, pp. 342–352. Springer, 2018.
[7] Igor V. Filippov and Marc C. Nicklaus. Optical structure recognition software to recover chemical information: Osra, an open source solution. *J. Chem. Inf. Model.*, 49(3):740–743, 2009. URL [https://doi.org/10.1021/ci800067r](https://doi.org/10.1021/ci800067r).

[8] Lin Gao, Weidong Song, Jiguang Dai, and Yang Chen. Road extraction from high-resolution remote sensing imagery using refined deep residual convolutional neural network. *Remote Sensing*, 11(5):552, 2019.

[9] Anna Gaulton, Anne Hersey, Michal Nowotka, A Patricia Bento, Jon Chambers, David Mendez, Prudence Muttoo, Francis Atkinson, Louise J Bellis, Elena Cibrán-Uhalte, et al. The chembl database in 2017. *Nucleic acids research*, 45(D1):D945–D954, 2017.

[10] Steven Gold and Anand Rangarajan. Graduated assignment graph matching. In *Proceedings of International Conference on Neural Networks (ICNN’96)*, volume 3, pp. 1474–1479. IEEE, 1996.

[11] Chi Han, Jiayuan Mao, Chuang Gan, Josh Tenenbaum, and Jiajun Wu. Visual concept-metaconcept learning. In *Advances in Neural Information Processing Systems*, pp. 5001–5012, 2019.

[12] Corentin Henry, Seyed Majid Azimi, and Nina Merkle. Road segmentation in sar satellite images with deep fully convolutional neural networks. *IEEE Geoscience and Remote Sensing Letters*, 15(12):1867–1871, 2018.

[13] Dakshina Ranjan Kisku, Ajita Rattani, Enrico Grosso, and Massimo Tistarelli. Face identification by sift-based complete graph topology. In *2007 IEEE workshop on automatic identification advanced technologies*, pp. 63–68. IEEE, 2007.

[14] Xiangnan Kong, Jiawei Zhang, and Philip S Yu. Inferring anchor links across multiple heterogeneous social networks. In *Proceedings of the 22nd ACM international conference on Information & Knowledge Management*, pp. 179–188, 2013.

[15] Wouter Marco Kouw and Marco Loog. A review of domain adaptation without target labels. *IEEE transactions on pattern analysis and machine intelligence*, 2019.

[16] Harold W Kuhn. The hungarian method for the assignment problem. *Naval research logistics quarterly*, 2(1-2):83–97, 1955.

[17] Francesco Locatello, Dirk Weissenborn, Thomas Unterthiner, Aravindh Mahendran, Georg Heigold, Jakob Uszkoreit, Alexey Dosovitskiy, and Thomas Kipf. Object-centric learning with slot attention. *arXiv preprint arXiv:2006.15055*, 2020.

[18] Noureddin M. Sadawi, Alan Sexton, and Volker Sorge. Chemical structure recognition: A rule based approach. *Proc. SPIE*, 8297:32–, 01 2012. URL [https://doi.org/10.1117/12.912185](https://doi.org/10.1117/12.912185).

[19] Jiayuan Mao, Chuang Gan, Pushmeet Kohli, Joshua B Tenenbaum, and Jiajun Wu. The neuro-symbolic concept learner: Interpreting scenes, words, and sentences from natural supervision. *arXiv preprint arXiv:1904.12584*, 2019.

[20] Joe R. McDaniel and Jason R. Balmuth. Kekule: Ocr-optical chemical (structure) recognition. *J. Chem. Inf. Comput. Sci.*, 32(4):373–378, 1992. URL [https://doi.org/10.1021/ci00008a018](https://doi.org/10.1021/ci00008a018).

[21] Martijn Oldenhof, Adam Arany, Yves Moreau, and Jaak Simm. Chemgrapher: optical graph recognition of chemical compounds by deep learning. *Journal of Chemical Information and Modeling*, 60(10):4506–4517, 2020.

[22] Jungkap Park, Gus Rosania, Kerby A Shedden, Mandee Nguyen, Naesung Lyu, and Kazuhiro Saitou. Automated extraction of chemical structure information from digital raster images. *Chem. Cent. J.*, 3:4, 03 2009. URL [https://doi.org/10.1186/1752-153X-3-4](https://doi.org/10.1186/1752-153X-3-4).

[23] Judea Pearl. Causal diagrams for empirical research. *Biometrika*, 82(4):669–688, 1995.
[24] Judea Pearl. Direct and indirect effects. In Proceedings of the Seventeenth conference on Uncertainty in artificial intelligence, pp. 411–420, 2001.

[25] Judea Pearl. Causality. Cambridge university press, 2009.

[26] Judea Pearl et al. Models, reasoning and inference. Cambridge, UK: Cambridge University Press, 2000.

[27] Óscar Pérez and Manuel Sánchez-Montañés. A new learning strategy for classification problems with different training and test distributions. In International Work-Conference on Artificial Neural Networks, pp. 178–185. Springer, 2007.

[28] Tyler Peryea, Daniel Katzel, Tongan Zhao, Noel Southall, and Dac-Trung Nguyen. Molvec: Open source library for chemical structure recognition. In ABSTRACTS OF PAPERS OF THE AMERICAN CHEMICAL SOCIETY, volume 258. AMER CHEMICAL SOC 1155 16TH ST, NW, WASHINGTON, DC 20036 USA, 2019.

[29] Alberto Sanfeliu and King-Sun Fu. A distance measure between attributed relational graphs for pattern recognition. IEEE transactions on systems, man, and cybernetics, (3):353–362, 1983.

[30] Francesc Serratosa. Fast computation of bipartite graph matching. Pattern Recognition Letters, 45:244–250, 2014.

[31] Francesc Serratosa. Computation of graph edit distance: reasoning about optimality and speed-up. Image and Vision Computing, 40:38–48, 2015.

[32] Francesc Serratosa. Speeding up fast bipartite graph matching through a new cost matrix. International Journal of Pattern Recognition and Artificial Intelligence, 29(02):1550010, 2015.

[33] Joshua Staker, Kyle Marshall, Robert Abel, and Carolyn M. McQuaw. Molecular Structure Extraction from Documents Using Deep Learning. J. Chem. Inf. Model., 59(3):1017–1029, March 2019. ISSN 1549-9596. doi: 10.1021/acs.jcim.8b00669. URL https://doi.org/10.1021/acs.jcim.8b00669

[34] Aniko T. Valko and A. Peter Johnson. Clide pro: The latest generation of clide, a tool for optical chemical structure recognition. J. Chem. Inf. Model., 49(4):780–787, 2009. URL https://doi.org/10.1021/ci800449t

[35] Alexander Vezhnevets, Vittorio Ferrari, and Joachim M Buhmann. Weakly supervised structured output learning for semantic segmentation. In 2012 IEEE conference on computer vision and pattern recognition, pp. 845–852. IEEE, 2012.

[36] David Weininger. SMILES, a chemical language and information system. 1. introduction to methodology and encoding rules. J. Chem. Inf. Comput. Sci., 28(1):31–36, 1988. URL https://pubs.acs.org/doi/abs/10.1021/ci00057a005

[37] Peter Willett, John M Barnard, and Geoffrey M Downs. Chemical similarity searching. . Chem. Inf. Comput. Sci., 38(6):983–996, 1998.

[38] Jia Xu, Alexander G Schwing, and Raquel Urtasun. Tell me what you see and i will show you where it is. In Proceedings of the IEEE conference on computer vision and pattern recognition, pp. 3190–3197, 2014.

[39] Jia Xu, Alexander G Schwing, and Raquel Urtasun. Learning to segment under various forms of weak supervision. In Proceedings of the IEEE conference on computer vision and pattern recognition, pp. 3781–3790, 2015.

[40] Zhe Xu, Shaoli Huang, Ya Zhang, and Dacheng Tao. Augmenting strong supervision using web data for fine-grained categorization. In Proceedings of the IEEE international conference on computer vision, pp. 2524–2532, 2015.

[41] Kexin Yi, Jiajun Wu, Chuang Gan, Antonio Torralba, Pushmeet Kohli, and Josh Tenenbaum. Neural-symbolic vqa: Disentangling reasoning from vision and language understanding. In Advances in neural information processing systems, pp. 1031–1042, 2018.
[42] Honghui Zhang, Jianxiong Xiao, and Long Quan. Supervised label transfer for semantic segmentation of street scenes. In European Conference on Computer Vision, pp. 561–574. Springer, 2010.

A APPENDIX

A.1 ARCHITECTURE SUMMARY OF GRAPH RECOGNITION TOOL

Every iteration of our method we need to train the graph recognition tool described in Oldenhof et al. [21]. This graph recognition tool is built using a combination of different convolutional neural networks. The first part is a semantic segmentation network to pixel-wise predict every atom, bond and charge type. The second part consists of three classification networks to classify every segment predicted by the semantic segmentation network. After the first step of the ChemGrapher model [21], the segmentation network, the predicted segments are processed so that for every segment the center of mass is calculated. These centers of mass would be the atom/bond/charge candidates to be classified by the classification networks.

| Layer   | Kernel | Nonlinearity | Padding | Dilation |
|---------|--------|--------------|---------|----------|
| conv1   | 3x3    | ReLU         | 1       | no dilation |
| conv2   | 3x3    | ReLU         | 2       | 2         |
| conv3   | 3x3    | ReLU         | 4       | 4         |
| conv4   | 3x3    | ReLU         | 8       | 8         |
| conv5   | 3x3    | ReLU         | 8       | 8         |
| conv6   | 3x3    | ReLU         | 4       | 4         |
| conv7   | 3x3    | ReLU         | 2       | 2         |
| conv8   | 3x3    | ReLU         | 1       | no dilation |
| last    | 1x1    | none         | no padding | no dilation |

Table 3: Summary of the layers of the segmentation network

| Layer   | Kernel | Nonlinearity | Padding | Dilation |
|---------|--------|--------------|---------|----------|
| depthconv1 | 3x3    | ReLU         | 1       | no dilation |
| conv2   | 3x3    | ReLU         | 2       | 2         |
| conv3   | 3x3    | ReLU         | 4       | 4         |
| conv4   | 3x3    | ReLU         | 8       | 8         |
| conv5   | 3x3    | ReLU         | 1       | no dilation |
| global maxpool | input size | None | no padding | no dilation |
| last    | 1x1    | None         | no padding | no dilation |

Table 4: Different layers in the classification network

A.2 TRAINING DETAILS FOR GRAPH RECOGNITION TOOL

Training details of the graph recognition tool for every iteration of our method are summarized in Table 5. The input images used for training of the different networks are a mix if images from source domain and upsampled rich labeled images from target domain. For pretraining of the ChemGrapher model only images from source domain were used. The training was performed using a compute node with 2 NVIDIA v100 GPUs with 32GB of memory.
Table 5: Training details for different networks

| Network       | #input images | #epochs | walltime | minibatch size | learning rate |
|---------------|---------------|---------|----------|----------------|---------------|
| Segm. network | 114K          | 5       | 24h      | 8              | 0.001         |
| Atom Clas.    | 12.4K         | 2       | 8h       | 16             | 0.001         |
| Charge Clas.  | 12.4K         | 2       | 8h       | 16             | 0.001         |
| Bond Clas.    | 4.4K          | 2       | 4h       | 64             | 0.001         |

A.3 Computational cost per rich-labeling iteration

In the following Table 6 the computational cost for 1 rich-labeling iteration is summarized including all steps: (re)training, predicting and graph aligning rich-labeling.

Table 6: Computational costs per rich-labeling iteration

| Hardware       | Training          | Predict          | Graph Aligning   |
|----------------|-------------------|------------------|------------------|
| Dataset        | 2 NVIDIA v100 GPUs| 1 NVIDIA v100 GPU| Intel Xeon Gold 6240 2.6Ghz |
| #datapoints    | see Table 5        | 4,000            | 4,000            |
| Walltime       | ∼44h (details Table 5) | ∼2h              | ∼40min           |

A.4 Examples of cases where graph alignment fails

We would like to showcase some examples where the constrained (max 2 node substitutions or max 1 edge substitution) graph alignment fails. At the same time it is important to note that our proposed domain adaptation method is an iterative method, so if a graph alignment fails in a previous iteration it could succeed in a next one when the new model makes a new graph prediction closer to the true graph.
Figure 7: Example 1: It is clear that to align the graph prediction $W'$ with the true graph $W$ more than 2 node substitutions are needed. So no rich labeling is possible for this example in this iteration.
Figure 8: Example 2: It is clear that alignment of the graph prediction $W'$ with the true graph $W$ can not be solved with only substitutions.

Figure 9: Example 3: It is clear that alignment of the graph prediction $W'$ with the true graph $W$ can not be solved with only substitutions.