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Molecule Edit Graph Attention Network: Modeling Chemical Reactions as Sequences of Graph Edits

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Abstract

One of the key challenges in automated chemical synthesis planning is to propose diverse and reliable reactions. A common approach is to generate reactions using reaction templates, which represent a reaction as a fixed graph transformation. This enables accurate and interpretable predictions but can suffer from limited diversity. On the other hand, template-free methods increase diversity but can be prone to making trivial mistakes. Inspired by the efficacy of reaction templates, we propose Molecule Edit Graph Attention Network (MEGAN), a template-free model that encodes reaction as a sequence of graph edits. Our model achieves state-of-the-art results on a standard retrosynthesis benchmark without any manual rule encoding.

1. Introduction

Chemical synthesis planning is a demanding task due to the substantial size and complexity of the reaction space.
Computer-aided methods are a promising approach to automate this process (Corey & Wipke, 1969; Segler et al., 2018; Coley et al., 2018a; Lee et al., 2019).

Generating reaction candidates is an essential part of automated synthesis planning. The accuracy with which we are able to predict reaction outcomes remains a key roadblock in wider applicability of computer-aided methods (Davies, 2019).

Many of the recent approaches to reaction generation use *reaction templates*, which are encoded graph transformation rules that enable generating reactions by applying them to the input molecule. Such methods are highly interpretable and achieve strong performance (Dai et al., 2020). However, template-based methods have some disadvantages. Perhaps

the most pressing one is that due to the computational limits they require modeling reactions using a relatively small number of templates. This necessarily limits the size of the chemical reaction space accessible by such methods.

The shortcomings of template-based methods have been addressed with the development of template-free models for reaction generation. In particular, modeling reaction generation as a machine translation task has lead to promising results (Liu et al., 2017). This and related methods are still however often outperformed by template-based methods (Dai et al., 2020), and have been also argued to be prone to making trivial mistakes (Molga et al., 2019).

Arguably, a more natural approach to reaction generation is to represent reaction as a sequence of graph edits. Bradshaw et al. (2018) models reaction as a sequence of bond removals and additions. However, their approach is limited to a certain subset of the chemical reaction space (reactions with linear chain topology) and forward synthesis (predicting product of a reaction). Do et al. (2019) models reaction as a set of operation on atom pairs. Similarly, their method cannot be readily applied to retrosynthesis (predicting substrates based on the product) due to the lack of support for atom addition.

In this work, we present the Molecule Edit Graph Attention Network (MEGAN). We propose an encoder-decoder model that generates a reaction as a sequence of graph edits. We include atom addition and bond removal in the action space to apply the model to the retrosynthesis task, where we achieve state-of-the-art results. Specifically, our main contributions are as follows:

- We propose a novel graph encoder-decoder architecture for reaction generation.
- We achieve state-of-the-art results in retrosynthesis.
- We demonstrate the feasibility of representing reaction as a sequence of graph edits for retrosynthesis, which is a promising step towards more interpretable neural models for reaction generation.

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Molecule Edit Graph Attention Network



Figure 1. Retrosynthesis prediction generated by MEGAN. The model modifies the target molecule by sequentially executing actions on the molecular graph until it decides to stop.

2. Molecule Edit Graph Attention Network

Graph Convolutional Networks (GCN) have been successfully applied to a variety of tasks in computational chemistry (Duvenaud et al., 2015; Xie & Grossman, 2018; You et al., 2018a; Coley et al., 2018b). Our Molecule Edit Graph Attention Network (MEGAN) is an encoder-decoder architecture based on GCN that is able to predict a sequential series of actions on atoms and bonds of a chemical compound. The key innovation is to combine an effective architecture and training procedure, which enables us to extend ideas from Bradshaw et al. (2018); Do et al. (2019) to retrosynthesis and achieve state of the art performance. We begin by describing the input and the output representation. Next, we introduce the overall architecture.

2.1. Input and output representation

Reaction as a sequence of edits We reformulate retrosynthesis as predicting a series of actions on the graph of the product molecule that produce the reactants. We define the following graph actions:

- Edit atom properties (*EditAtom*)
- Edit bond between two atoms (EditBond)
- Add new atom to the graph (AddAtom)
- Add new benzene ring to the graph (AddBenzene)
- Stop generation (*Stop*)

EditAtom changes properties of atoms, such as the formal charge, chirality or aromaticity. *EditBond* adds, edits or deletes a bond between two atoms. *AddAtom* adds a new atom of a specified type as a neighbor of another atom already existing in the graph, with a specified bond type. *AddBenzene* optimizes atom addition by appending a complete benzene ring to a selected carbon atom. *Stop* action

indicates the end of the generation process. We use atom mapping information to define an order on actions, which we describe in the experimental section. We describe the possible actions in details in the Supplement.

Molecule representation MEGAN takes as input a molecular graph, which is represented by labeled node vectors and a labeled adjacency matrix. More specifically, the input consists of a matrix of features $H^{OH} \in \mathbb{Z}_{\geq 0}^{n \times h_{OH}}$ and an adjacency matrix $A^{OH} \in \mathbb{Z}_{\geq 0}^{n \times n \times a_{OH}}$, where *n* is the number of nodes in a graph and h_{OH} and a_{OH} are sizes of concatenated one-hot vectors of atom and bond features. Hydrogen atoms are removed from the graph, as for heavy atoms the number of neighboring hydrogen atoms can be deduced implicitly or marked explicitly by a special atom feature if needed. We selected a minimal set of atom and bond features that allows for exact reconstruction of SMILES of all products and reactants from the development set. We describe the featurization in details in the Supplement.

2.2. Model architecture

Input embedding Input features are embedded using linear layers $f_{emb} : \mathbb{R}^{h_{OH}} \to \mathbb{R}^{h}$ and $g_{emb} : \mathbb{R}^{a_{OH}} \to \mathbb{R}^{a}$:

$$\mathbb{R}^{n \times h} \ni H^0 = f_{emb}(H^{OH}) \tag{1}$$

$$\mathbb{R}^{n \times n \times a} \ni A = g_{emb}(A^{OH}) \tag{2}$$

GCN-att layer The basic building block of the network is an attention-based GCN layer named *GCN-att*. We enhance the GCN layer from Veličković et al. (2017) by adding bond features as input information for computing the attention values. Let $H^t \in \mathbb{R}^{n \times h}$ denote input node features for the *t*-th GCN-att layer and $N(i) \subset \mathbb{Z}_{\geq 0}$ denote set of indices of neighbors of node at index *i* (where $i \in N(i)$). We calculate new node features $H^{t+1} \in \mathbb{R}^{n \times h}$ as follows:

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$$\mathbb{R}^d \ni H_i^{t\prime} = \sigma_r(f_{att}^t(H_i^t)) \tag{3}$$

$$\mathbb{R}^{2d+a} \ni B_{ij}^t = H_i^{t\prime} \parallel H_j^{t\prime} \parallel A_{i,j} \tag{4}$$

$$\mathbb{R}^K \ni C_{ij}^t = f_{att'}^t(B_{ij}^t) \tag{5}$$

$$\mathbb{R}^{h} \ni G_{ik}^{t} = \sum_{j \in N(i)} \frac{\exp C_{ijk}^{t}}{\sum_{i \in N(i)} \exp C_{ilk}^{t}} H_{j}^{t} \qquad (6)$$

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 $\mathbb{R}^h \ni H_i^{t+1} = \underset{1 \le k \le K}{\parallel} \sigma_r(f_k^t(G_i^t)) \tag{7}$

where σ_r denotes *relu* activation function, $\|$ indicates vector concatenation, $K \in \mathbb{N}_+$ is the number of attention heads and $f_{att} \colon \mathbb{R}^h \to \mathbb{R}^d$, $f_{att'} \colon \mathbb{R}^{2d+a} \to \mathbb{R}^K$ and $f \colon \mathbb{R}^h \to \mathbb{R}^{h/K}$ are standard linear layers. Numbers h, a, d and K are hyperparameters of the model. We require that h is divisible by K. We use the same hyperparameter values for all GCNatt layers in the model, but do not share their weights.

129 Supernode A single pass through a GCN-att layer trans-130 fers information only between neighboring atoms. This can 131 potentially hinder the ability to learn graph-level features, 132 such as coexistence of functional groups in different parts 133 of a compound. To mitigate this, we introduce an additional 134 node named supernode (Li et al., 2017), which is connected 135 to all atoms in the graph with a special SUPERNODE bond 136 type. Supernode is particularly useful for passing informa-137 tion between connected components of a graph after it was 138 split by deleting a bond. 139

Overall architecture The model consists of two parts: the 140 encoder, which is invoked only once per reaction generation 141 and the *decoder*, which is sequentially invoked to generate 142 actions (Figure 2). At each generation step, features are 143 embedded using layers f_{emb} and g_{emb} . The encoder has 144 N_e stacked GCN-att layers, which are invoked at the first 145 generation step after the embedding layers. The decoder has N_d stacked GCN-att layers followed by final linear layers 147 that output action probabilites for atom and bond actions 148 as follows. Let $m = N_e + N_d$ for the first generation step 149 and $m = N_d$ for the other generation steps. The logits L^a 150 and L^b for atom actions Act_a and bond actions Act_b are 151 calculated as follows: 152

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$$\mathbb{R}^d \ni F_i = \sigma_r(g_{atom}(H_i^m)) \tag{8}$$

$$\mathbb{R}^{|Act_a|} \ni L_i^a = g'_{atom}(F_i) \tag{9}$$

$$\mathbb{R}^d \ni J_i = \sigma_r(g_{bond}(H_i^m)) \tag{10}$$

$$\mathbb{R}^{d+a} \ni J'_{ij} = \sigma_r(J_i + J_j \parallel A_{i,j}) \tag{11}$$

$$\mathbb{R}^{|Act_b|} \ni L^b_{ij} = g'_{bond}(J'_{ij}) \tag{12}$$

We reuse the hyperparameter value d for simplicity. To acquire the final action probabilities, we apply softmax ac-

tivation function to concatenated vectors of logits of all possible atom actions Act_a and possible bond actions Act_b . To decide which actions are legal, we use the following rules:

- Stop action can be predicted only by the supernode
- All other atom actions can be predicted by all nodes except the supernode.
- Bond actions can be predicted for indices *i* and *j*, where *i* < *j* and nodes at *i* and *j* are atoms

Retaining generation state Finally, we want our model to be *stateful*, that is to be able to take advantage of the information about the previous generation steps to predict the next action. We achieve this by storing the output H_s^m of the last GCN-att layer at the generation step s and merging it with the embedded input H_{s+1}^0 at step s+1: $\mathbb{R}^h \ni H_{s+1}^0 := \max(H_{s+1}^0, H_s^m)$, where max is the piecewise maximum of vectors. In H_s^m , we zero-pad features for any node that was added to the graph at step s.

3. Experiments

We run experiments on the standard retrosynthesis benchmark USPTO-50k (Lowe, 2012; Schneider et al., 2016). We begin by introducing the experimental setting.

3.1. Experimental setting

Tasks We evaluate the models on retrosynthesis. The goal in retrosynthesis is to predict the set of reactants based on the product of a reaction. The accuracy is measured by comparing the SMILES (Weininger, 1988) representation of the generated reactants to the SMILES representation of the ground truth reactants. Before the comparison, we remove any mapping information from the SMILES strings and canonicalize them using RdKit (Landrum). We use top K accuracy computed on the reactions from the test set as the main evaluation metric.

Data We use the USPTO-50k data set of approximately 50000 reactions, which was collected by Lowe (2012) and classified into 10 reaction types by Schneider et al. (2016). We use the same processed version of the data set as Coley et al. (2017), where each reaction consists of a single product molecule and a set of one or more reactants, with corresponding atoms between the reactants and the product mapped. Following other studies, we assign each reaction randomly to one of the training/validation/test sets with respective probabilities of 80%/10%/10%.

Gradient-based training of MEGAN In contrast to Do et al. (2019) who use reinforcement learning to train their model, we back-propagate directly through the maximum

165 likelihood objective to train MEGAN. This is nontrivial, as computing the gradient of the likelihood objective requires 167 defining a fixed ordering of actions (You et al., 2018b). To 168 solve this issue, You et al. (2018b) enumerates atoms using 169 breadth-first search. We adapt a similar idea to reaction gen-170 eration. We use the mapping provided in the USPTO-50k 171 data set, which describes atom correspondence between the 172 product and the substrates, to predetermine an ordering of 173 actions. This provides supervision for each generation step 174 and thus enables us to compute the gradient. We provide 175 the remaining details in the Supplement.

176 Other training and evaluation details of MEGAN For 177 training, we use batch size of 4 reactions. We use 178 Adam (Kingma & Ba, 2014) with the initial learning rate of 179 0.0001. We use warm-up, increasing the learning rate from 180 0 to 0.0001 over the first 10000 training steps. For efficiency, 181 we compute the validation loss on a subset of 2500 valida-182 tion samples after each 10000 training samples. We multiply 183 the learning rate by 0.1 if the estimated validation loss has 184 not decreased for more than 4 such validation checks. We 185 stop the training after the estimated validation loss has not 186 decreased for more than 8 validation checks. We adapt the 187 hyperparameters based on the validation loss. The final 188 hyperparameter values are described in the Supplement. 189

Following other studies, we run two variants of training:
one with unknown reaction type and one for which reaction
type is given as a prior by an additional embedding layer.
For both runs, we use the same model architecture and the
same training setup. The training takes approximately 8
hours on a single Nvidia Tesla V100 GPU for both variants.

196 We use beam search (Graves, 2012) on output probabilities 197 of actions to generate multiple ranked candidates for each 198 product. We set the maximum number of steps to 16 and the 199 beam width to 50, as it is the largest K for which accuracy 200 was reported for the baseline models. Running predictions with the beam width of 50 on the test set of USPTO-50k 202 (5155 test products) takes about 45 minutes on a single 203 Nvidia Tesla V100 GPU. 204

Baselines We compare performance of MEGAN with sev-206 eral template-free and template-based models, including current state-of-the-art methods. Seq2seq (Liu et al., 2017) 208 and Transformer (Karpov et al., 2019) are both template-209 free methods based on machine translation models applied 210 on SMILES strings. G2G (Shi et al., 2020) (concurrent 211 work) is also a template-free model based on modifying 212 molecular graphs, with a separate module for predicting 213 reaction center. Retrosim (Coley et al., 2017) uses reaction 214 fingerprint to select template based on similar reactions in 215 the data set. Neuralsym (Segler & Waller, 2017) uses a 216 multi-linear percceptron to rank templates. GLN (Dai et al., 217 2020) employs a graph model that assesses when rules from 218 templates should be applied.

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Table 1. Top-k test accuracy on the USPTO-50k data set. Results of other methods taken from Dai et al. (2020) and Shi et al. (2020)

METHODA	Тор-к accuracy %						
Methods	1	3	5	10	20	50	
	REACT	TION TY	PE UNKI	NOWN			
TRANS	37.9	57.3	62.7	/	/	/	
Retrosim	37.3	54.7	63.3	74.1	82.0	85.3	
NEURALSYM	44.4	65.3	72.4	78.9	82.2	83.1	
G2Gs	48.9	67.6	72.5	75.5	/	/	
GLN	52.5	69.0	75.6	83.7	89.0	92.4	
MEGAN	47.6	71.2	79.5	87.0	91.4	94.2	
R	EACTIO	N TYPE	GIVEN A	AS PRIO	R		
SEQ2SEQ	37.4	52.4	57.0	61.7	65.9	70.7	
RETROSIM	52.9	73.8	81.2	88.1	91.8	92.9	
NEURALSYM	55.3	76.0	81.4	85.1	86.5	86.9	
G2Gs	61.0	81.3	86.0	88.7	/	/	
GLN	64.2	79.1	85.2	90.0	92.3	93.2	
MEGAN	61.6	83.7	89.2	93.2	95.3	96.6	

3.2. Results

Table 1 reports results on the USPTO-50k benchmark in a variant with and without reaction type information. For the top ranked prediction (K = 1), MEGAN is surpassed only by GLN when reaction type is given and by GLN and G2Gs when reaction type is unknown. For K > 1, our model achieves state-of-the-art accuracy, outperforming all baselines in both settings.

We hypothesize that the advantage of MEGAN for K > 1 stems largely from the fact that MEGAN generates reaction as a sequence of edits. This might help to efficiently search through different plausible reaction centers, hence covering a more diverse subset of the reaction space. It can also enable MEGAN to achieve high coverage of the reaction space, which is indicated by Top 50 accuracy of 94.2% when reaction type is unknown and 96.6% when reaction type is provided.

4. Conclusions

In this work, we presented the Molecule Edit Graph Attention Network (MEGAN), a template-free model that achieves state-of-the-art performance on retrosynthesis. The key idea is to generate reaction as a sequence of edits. We hypothesize that this enables generating more diverse predictions by a more efficient search through the space of plausible reactions.

To train our model efficiently, we used heuristically computed reaction mapping to define an order of actions. An interesting topic for the future would be to combine gradientbased training with reinforcement learning to reduce the reliance of MEGAN on mapping.

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330 5. Supplementary Material

5.1. Featurization

333 We featurize atoms and bonds with one-hot encoded vectors 334 of features calculated using RdKit (Landrum). We select 335 features that allow for correct reconstruction of all products and substrates from the USPTO-50k development set. 337 In Tables 2 and 3 we present all used features and their 338 possible values. We concatenate one-hot feature vectors 339 to gain the final input representation of atoms and bonds $H^{OH} \in \mathbb{Z}_{\geq 0}^{n \times h_{OH}}$ and $A^{OH} \in \mathbb{Z}_{\geq 0}^{n \times n \times a_{OH}}$. During 340 341 evaluation, if an unknown feature value is seen (for instance, 342 BOND TYPE=QUADRUPLE), we set the one-hot vector for 343 this feature to zeros. For supernodes, all one-hot feature 344 vectors are set to zeros, apart from vectors for the features 345 IS SUPERNODE and BOND TYPE. We connect each atom 346 with itself with a special bond of type SELF. For non-347 neighboring atoms at i and j we set $A_{ij}^{OH} = \vec{0}$.

For both atoms and bonds, we add a special IS EDITED feature that marks all bonds and atoms that have been modified by actions. This aims to help the decoder to focus on these atoms and bonds, as they are the most probable candidates for the next generation steps.

5.2. Graph edit actions

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In Table 6 we show all possible graph actions. The actions
were found during the generation of the training samples,
which we describe in the next paragraph. The actions have
different sets of parameters, depending on the action type.

For EditAtom, the action parameters are the atom properties 361 that are changed by the action. Note that a single EditAtom 362 action sets all these properties to the specified values. For 363 instance, action number 1, when executed on an atom, sets its formal charge to 0, chiral tag to None, number of explicit Hydrogen atoms to 1 and marks it as aromatic. EditBond acts similarly to *EditAtom* but edits properties of a bond 367 instead of an atom. AddAtom adds a new atom with specified features, connected to an existing atom with a bond with 369 specified features. AddBenzene appends a benzene ring 370 to a specified carbon atom and has no parameters. Stop 371 terminates reaction generation and also has no parameters. 372

BondEdit actions are bond actions, that is they are predicted for a pair of atoms. All other types of actions are atom actions and are predicted for a single atom in the graph.

5.3. Generating training samples

For each reaction from the development set, we generate
training samples by finding actions that lead from the target
molecule to the reactants. At first, we remap the reaction
to a canonical form so that the atom map numbers correspond to to the order of atoms in the canonical SMILES of

the compounds. We construct the target graph T and the reactants graph R, using features from Tables 2 and 3. We also initialize the stack of edited atoms $S = \{\}$, which is used for drawing candidate atoms for actions, prioritizing depth-first generation. At each step, we execute the first possible action for the concatenated lists of candidate atoms $S \parallel M(T)$, where M(T) is the list of all atoms from the target ordered by their mapping numbers. After each step, we push the modified atoms to the stack S. If there is more than one possible action to perform on a selected atom i or atom pair i, j, we use the following priority of actions Π :

- Action that deletes the existing bond between i, j
- Any action that adds a bond between i, j
- Any action that edits the existing bond between i, j
- Any action that edits the existing atom *i*
- Any action that adds a benzene ring to i
- Any action that adds an atom to i

where bond deletion has the highest priority. The aim is to prioritize actions that are usually the hardest (such as deleting a bond, which usually means finding the reaction center) or logically follow each other (editing a neighboring bond and/or atom often follows bond deletion or addition). The training reaction generation is described in Algorithm 1.

5.4. Hyperparameter search

Table 4 shows the hyperparameter values used for the final models. We found them heuristically by observing the validation loss when training with unknown reaction type. We tried increasing N_e to 8, as well as K to 12 and h to 1440 but we did not see a decrease in validation error. We also wanted the decoder to be as small as possible, as it is invoked multiple times during the reaction generation. We found that $N_d = 2$ is sufficient to achieve the best accuracy and increasing it lead to faster overfitting. The total number of learnable parameters in the model is approximately 9.8 million.

NAME	POSSIBLE VALUES		
Is supernode	YES, NO	2	
ATOMIC NUM-	5, 6, 7, 8, 9, 12, 14, 15, 16,	16	
BER	17, 29, 30, 34, 35, 50, 53		
FORMAL	-1, 0, 1	3	
CHARGE			
CHIRAL TAG	None, @, @@	3	
NUMBER OF	0, 1, 2, 4	4	
EXPLICIT HS			
IS AROMATIC	YES, NO	2	
Is edited	YES, NO	2	
TOTAL		32	

Table 3. Features of bonds							
NAME	POSSIBLE VALUES	Dim					
BOND TYPE	Supernode, Self, Single, Double, Triple, Aromatic	6					
BOND STEREO	None, Z, E	3					
IS EDITED	YES, NO	2					
TOTAL		11					

Table 4. Final model hyperparameter values								
h 1024	a 128	d 128	$\left \begin{array}{c}K\\8\end{array}\right $	$\begin{vmatrix} N_e \\ 6 \end{vmatrix}$	$\begin{vmatrix} N_d \\ 2 \end{vmatrix}$	TOTAL PARAMS ~9.8MIL		

Table 5.	USPTO-50k	data set	information

# TRAIN	# VALID	# TEST	# TOTAL
39662	5199	5155	50016

Algorithm 1 Generating training samples for a reaction **Input:** target graph T, reactants graph RInitialize $S \coloneqq \{\}$ (empty list) Let $M(T) = \{$ all atoms from T ordered by mapping $\}$ marker: while $T \neq R$ do for i in $S \parallel M(T)$ do for P in Π do for a in P do if a is an atom action then if applying a to i leads to R then generate training sample (T, a)apply action a to atom i in T $S \coloneqq \{i\} \parallel S$ go to marker end if else if a is a bond action then for j in $S \parallel M(T)$ do if applying a to i, j leads to R then generate training sample (T, a)apply action a to atoms i, j in T $S \coloneqq \{j, i\} \parallel S$ go to marker end if end for end if end for end for end for end while

	FORMAL C	CHARGE	CHIRAL TAG	NUM	of explicit H	IS IS A	IS AROMATIC	
	0		None		1		YES	
	+1		NONE		0		No	
	0		@@		1		No	
	0		None		0		No	
EditAtom	0		@		1		No	
	-1		None		0		No	
	0		None		0		YES	
	0		@		0		No	
	0		@@		0		No	
	-1		None		0		YES	
	0	D	None		1		No	
		BOND TY			BO	ND STEREO		
		NONE (DELETH				/ None		
		Single Triple				NONE		
EditBond		DOUBL				NONE		
		DOUBL				Z		
		DOUBL				E		
		AROMAT				NONE		
	ATOMIC	Formal	CHIRAL	NUMBER	Is	Bond	Bond	
	NUM	CHARGE	TAG	OF EXP HS	AROMATIC	TYPE	STEREO	
	35	0	NONE	0	No	SINGLE	NONE	
	6	0	NONE	0	No	SINGLE	None	
	8	0	NONE	0	No	DOUBLE	None	
	6	0	NONE	0	YES	SINGLE	NONE	
	8	0	NONE	0	No	SINGLE	NONE	
	17	0	NONE	0	No	SINGLE	NONE	
	7	0	NONE	0	No	SINGLE	NONE	
	16	0	NONE	0	No	SINGLE	NONE	
	53	0	NONE	0	No	SINGLE	None	
	8	-1	NONE	0	No No	SINGLE	NONE	
	5 9	0 0	None None	0	No	SINGLE	NONE	
	15		NONE	0 0	No No	SINGLE	None None	
	13 7	+1 +1	NONE	0	NO	Single Single	NONE	
	50	+1 0	NONE	0	NO	SINGLE	NONE	
	14	0	NONE	0	NO	SINGLE	NONE	
AddAtom	7	+1	NONE	0	No	DOUBLE	NONE	
	, 7	-1	NONE	0	No	DOUBLE	NONE	
	29	0	NONE	ů 0	No	SINGLE	NONE	
	16	+1	NONE	ů 0	No	SINGLE	NONE	
	12	+1	NONE	Ő	No	SINGLE	NONE	
	15	0	NONE	0	No	SINGLE	NONE	
	6	0	@@	1	No	SINGLE	NONE	
	30	+1	NONE	0	No	SINGLE	None	
	30	0	None	0	No	SINGLE	None	
	6	0	NONE	0	No	DOUBLE	NONE	
	15	0	None	0	No	DOUBLE	None	
	16	0	NONE	0	No	DOUBLE	NONE	
	7	0	NONE	0	No	DOUBLE	NONE	
	12	0	NONE	0	No	SINGLE	NONE	
	6	0	@@	0	No	SINGLE	NONE	
	6	0	@	1	No	SINGLE	None	
	6	0	NONE	0	No	DOUBLE	E	
4.110	6	0	None	0	YES	AROMATIC	None	
AddBenzene				/				
Stop				/				