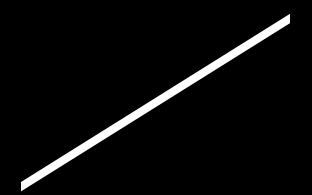
Learning from literatureextracted synthesis actions for organic synthesis





AMLD EPFL 30 March 2022

IBM Research

Data and chemical reactions

- Chemists have been doing reactions in roughly the same way for **decades**
- Set of standard lab operations
- Millions of reactions reported in the literature

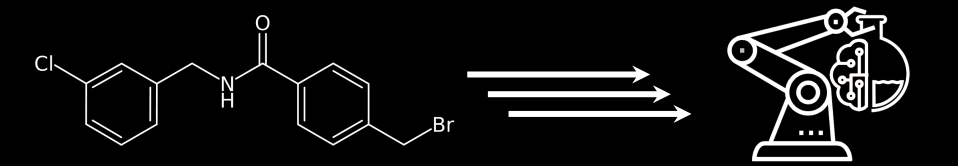




How can exploit this **data** to **accelerate discovery**?

- Assist chemists in synthesis planning
- ... and run the syntheses for them!

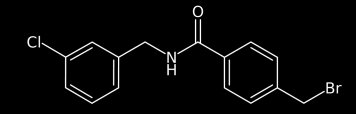
Data and chemical reactions



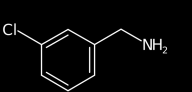
Target molecule

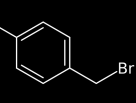
Synthesis execution

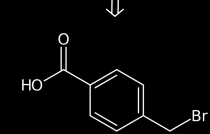
Step 1: retrosynthetic analysis



 \downarrow



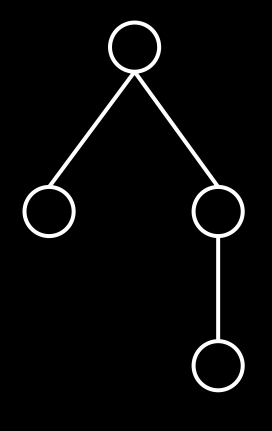




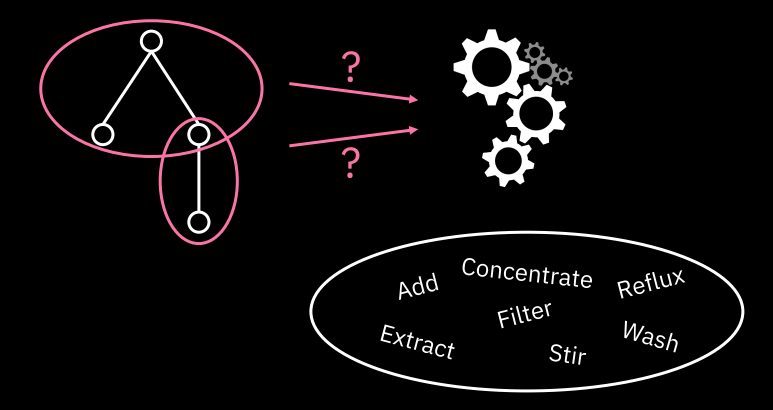
C

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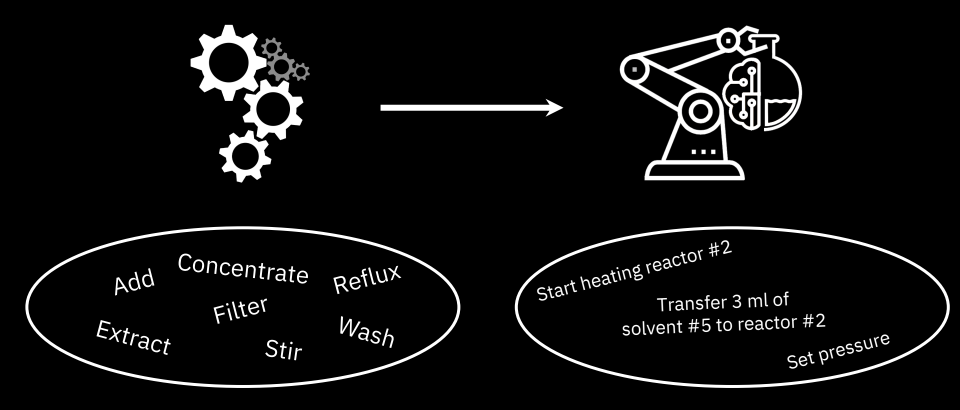
Retrosynthetic tree



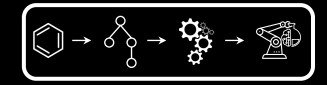
Step 2: experimental steps

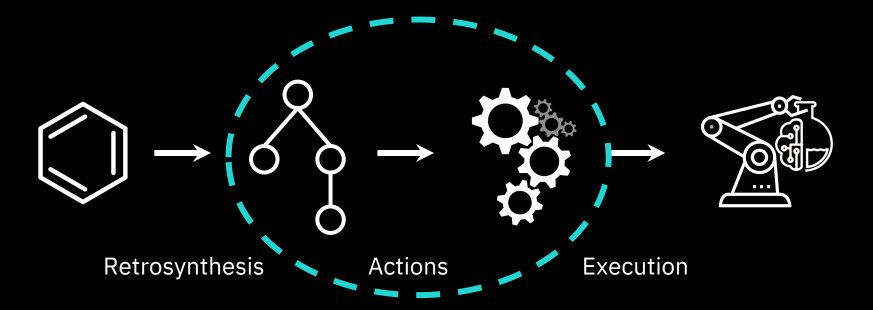


Step 3: Execution on robotic system

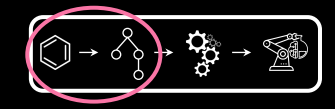


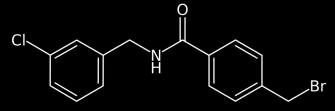
All together

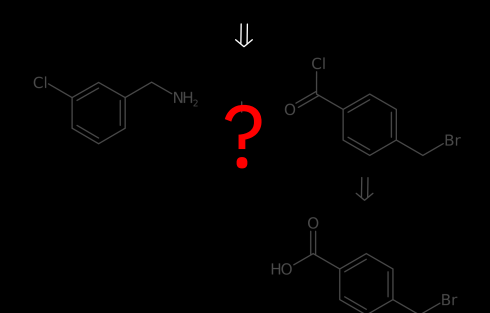




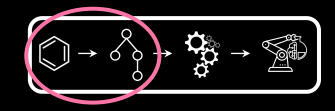
Retrosynthesis

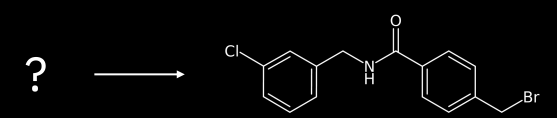


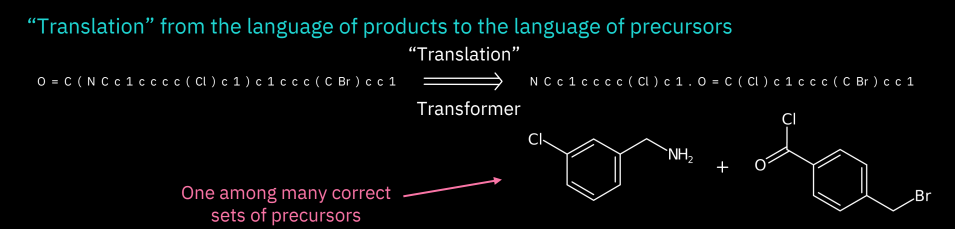








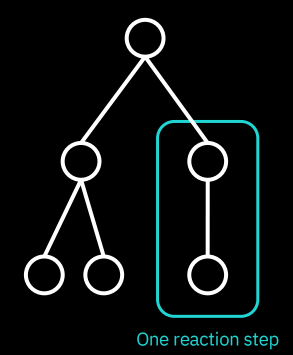


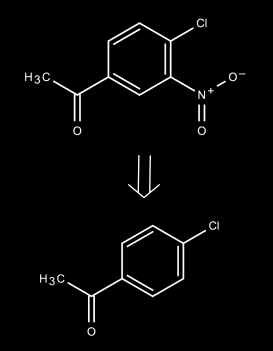


Schwaller, P.; Petraglia, R.; Zullo, V.; Nair, V. H.; Haeuselmann, R. A.; Pisoni, R.; Bekas, C.; Iuliano, A. & Laino, T., Chem. Sci., 2020, 11, 3316-3325.

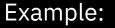
Synthesis actions

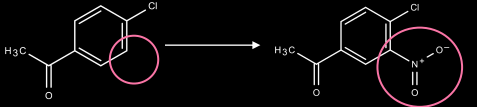


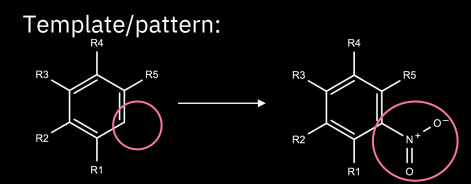




Synthesis actions



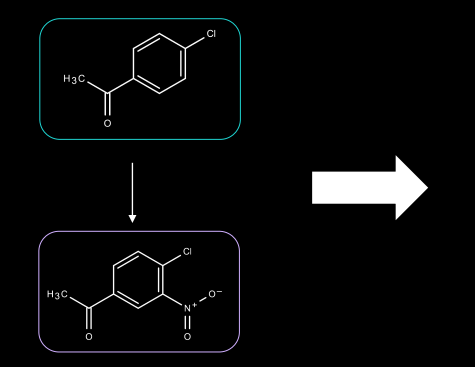




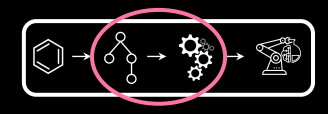


- -Same template but different synthesis actions!
- -Hard to predict
- –Ideally: ML model! "SMILES-to-actions"

Synthesis actions



C1=CC(C(=0)C)=CC=C1Cl>>C1=CC(C(=0)C)=CC([N+]([0-])=O)=C1Cl



Operation 1 Operation 2 Operation 3 Operation 4

SMILES-to-actions

– No dataset!

- Information is available indirectly
- First: extract actions from text
- "Paragraph-to-actions" model



Example procedure from a patent

A mixture of 1-(4-isopropyl-phenyl)-5-oxo-pyrrolidine-3-carboxylic acid ethyl ester obtained in step 2 (0.7 g, 2.65 mmol) and ethanol were cooled to 10-15° C. Sodium borohydride (0.25 g, 6.6 mmol) was added portion wise over a period of 20 min and the reaction mixture was stirred for 3.5 hrs at 20-25° C. The organic volatiles were evaporated and the residue was taken into brine solution (15 ml). The aqueous layer was extracted with ethyl acetate, dried over Na2SO4 and evaporated to obtain 4-hydroxymethyl-1-(4isopropyl-phenyl)-pyrrolidin-2-one as an off white solid (0.5 g, 81%).

Paragraph-to-actions: Action definition

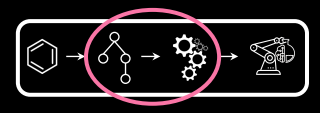
... Sodium borohydride (0.25 g, 6.6 mmol) was added portion wise over a period of 20 min and the reaction mixture was stirred for 3.5 hrs at 20-25° C ...





Paragraph-to-actions: Action definition

| Action name | Description | |
|---|---|-----------------------------|
| Add CellectLayer | Add a substance to the reactor Select aqueous or organic fraction(s) Evaporate the solvent (rotavap) | |
| Concentrate Crystallize Degas | Exaporate the solvent (rotavap) Re-crystallize a solid from a solvent Purge the reaction mixture with a gas | Action name |
| Dry <u>UnVacuum</u> DryWithMaterial Extract Filter | Dry a solid under vacuum Dry an organic solution with a desiccant Transfer compound into a different solvent Separate solid and liquid phases | Add |
| MakeSolution Microwave Partition PH | Mix several substances to generate a mixture or solution Heat the reaction mixture in a microwave apparatus Partition the reaction mixture by adding two immiscible solvents Change the pH of the reaction mixture | |
| PhaseSeparation Purify Quench Reflux | Separate the aqueous and organic phases Purification (chromatography) Stop reaction by adding a substance Reflux the reaction mixture | CollectLayer Concentrate |
| Reffux SetTemperature Sonicate Stir | Change the temperature of the reaction mixture Agitate the solution with sound waves Stir the reaction mixture for a specified duration | Crystallize Degas |
| Wait Wash Yield | Leave the reaction mixture to stand for a specified duration Wash (after filtration, or with immiscible solvent) Phony action, indicates the product of a reaction | |
| FollowOtherProcedure InvalidAction NoAction | The text refers to a procedure described elsewhere Unknown or unsupported action The text does not correspond to an actual action | |



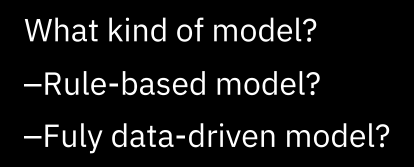
| ction name | Variable name | Variable type |
|------------|---------------|-------------------|
| d | material | chemical |
| | dropwise | boolean |
| | temperature | string (optional) |
| | atmosphere | string (optional) |
| | duration | string (optional) |
| llectLayer | layer | string |
| ncentrate | (none) | |
| ystallize | solvent | chemical |
| gas | gas | string (optional) |
| | duration | string (optional) |

Models for Paragraph-to-actions

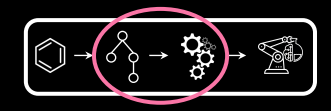
... Sodium borohydride (0.25 g, 6.6 mmol) was added portion wise over a period of 20 min and the reaction mixture was stirred for 3.5 hrs at 20-25° C ...

Add(name='Sodium borohydride', quantity=['0.25 g', '6.6 mmol'], duration='20 min')

Stir(temperature='20-25°C', duration='3.5 hrs')



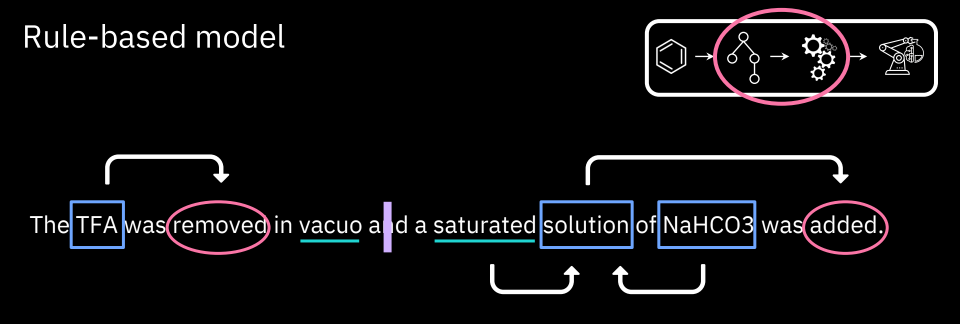
Models for Paragraph-to-actions



| Rule-based model | ML model | |
|----------------------------------|---------------------------------|--|
| Requires no training data | Requires training data | |
| Not very robust, hard to improve | Improve model by improving data | |

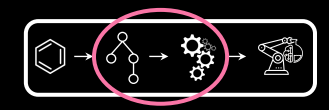
Let's combine both:

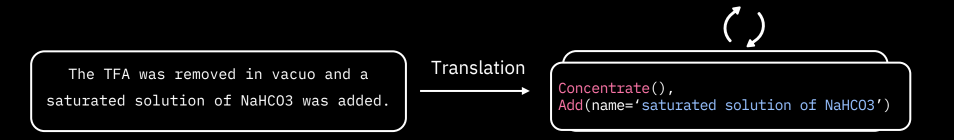
Rule-based training data for ML model



Generated actions for ~4M sentences

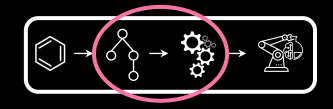
ML model





Transformer model, initial training on 4M samples

Rule-based vs ML model



Patent sentence: Diisopropylazodicarboxylate (0.05 ml, 0.302 mmol) was added to the reaction mixture followed by stirring for 3 hours at room temperature.

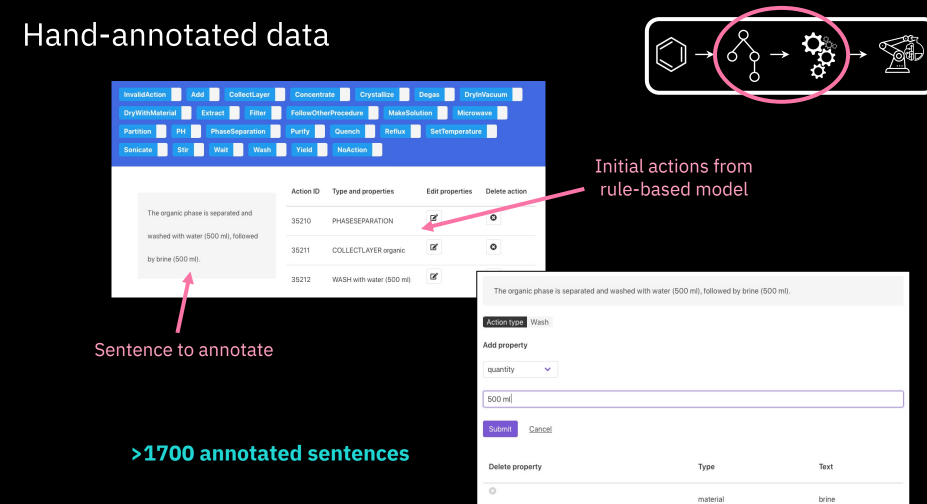
Rule-based model: ADD Diisopropylazodicarboxylate (0.05 ml, 0.302 mmol); STIR for 3 hours at room temperature. ML model: ADD Diisopropylazodicarboxylate (0.05 ml, 0.302 mmol); STIR for 3 hours at room temperature.

Patent sentence: The reaction mixture was concentrated in vacuo and water was added followed by enough hydrochloric acid (1 M) to acidify the solution.

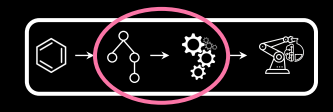
Rule-based model:CONCENTRATE; ADD water.ML model:CONCENTRATE; ADD water.Ground truth:CONCENTRATE; ADD water; PH with hydrochloric acid (1 M) to pH acidic.

Improving ML model:

- Training data size / quality
- Refine on human annotation

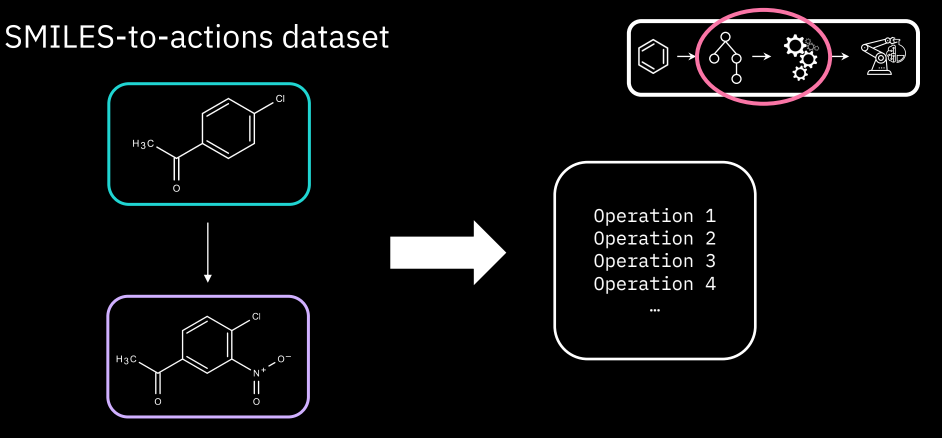


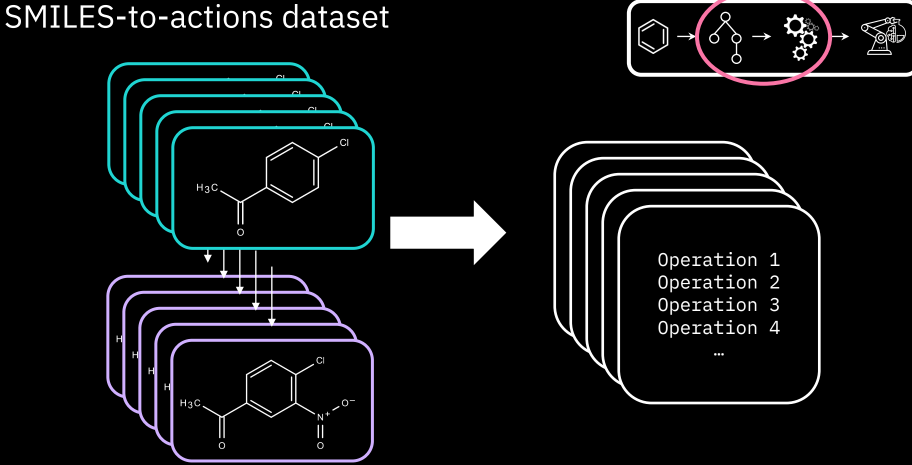
Results



| | 100% |
|------------------------------|----------|
| Model | accuracy |
| Combined rule-based model | 21.9 |
| Pretrained translation model | 24.7 |
| Model without pretraining | 37.8 |
| Refined translation model | 60.8 |

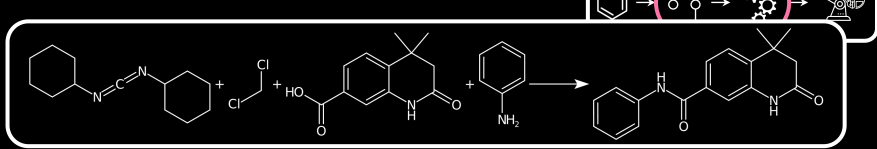
Vaucher, A. C.; Zipoli, F.; Geluykens, J.; Nair, V. H.; Schwaller, P.; Laino, T., *Nat. Commun.* **2020**, *11*, 3601.





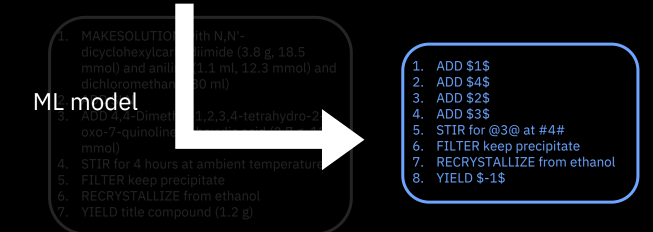
Vaucher, A. C.; Schwaller, P.; Geluykens, J.; Nair, V. H.; Iuliano, A.; Laino, T., *Nat. Commun.* **2021**, *21*, 2573. Alain Vaucher / IBM Research Europe / March 30, 2022

SMILES-to-actions



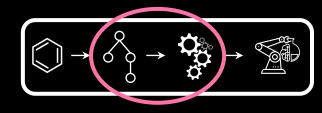
 $\texttt{C}(\texttt{=NC1CCCCC1})\texttt{=NC1CCCCC1} \ . \ \texttt{ClCl} \ . \ \texttt{CC1}(\texttt{C})\texttt{CC}(\texttt{=0})\texttt{Nc2cc}(\texttt{C}(\texttt{=0})\texttt{0})\texttt{ccc21} \ . \ \texttt{Nc1ccccc1} >> \ \texttt{CC1}(\texttt{C})\texttt{CC}(\texttt{=0})\texttt{Nc3ccccc3}\texttt{ccc21}$

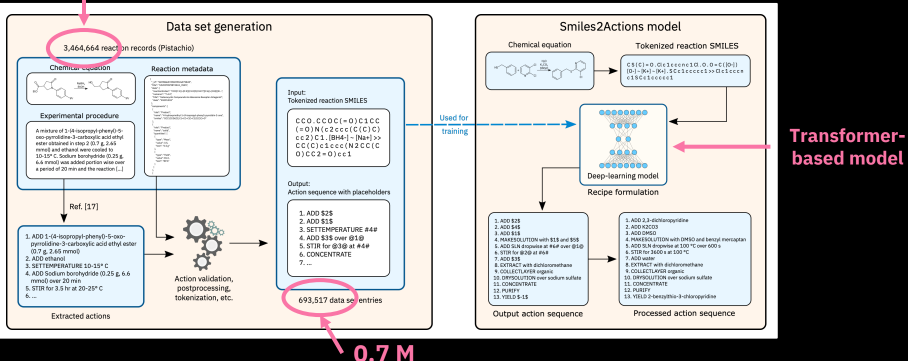
2.7 g (12.3 mmol) 4,4-Dimethyl-1,2,3,4tetrahydro-2-oxo-7-quinolinecarboxylic acid were added to a solution of 3.8 g (18.5 mmol) N,N'dicyclohexylcarbodiimide and 1.1 ml (12.3 mmol) aniline in 80 ml dichloromethane. The reaction mixture was stirred for 4 hours at ambient temperature and the precipitate was filtered off with suction and recrystallised from ethanol. There was obtained 1.2 g of the title compound; m.p. 249-251° C.



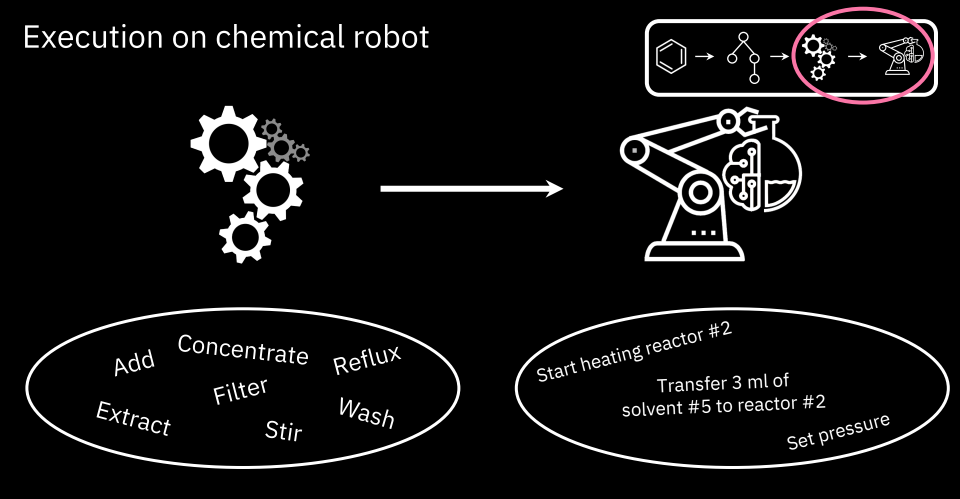
SMILES-to-actions

3.5 M

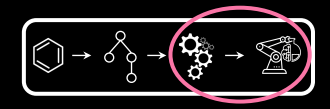




Vaucher, A. C.; Schwaller, P.; Geluykens, J.; Nair, V. H.; Iuliano, A.; Laino, T., *Nαt. Commun.* **2021**, *21*, 2573.



Execution on chemical robot



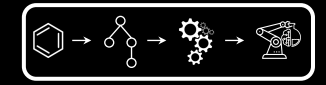
Cloud-based setup for autonomous synthesis

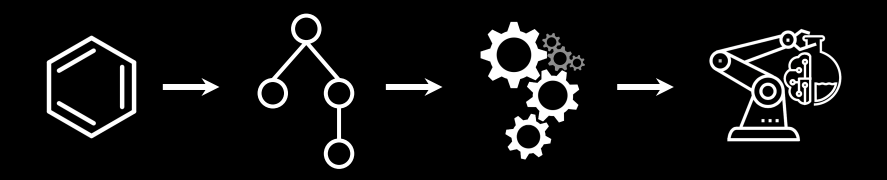


Hardware @ IBM Research Zurich

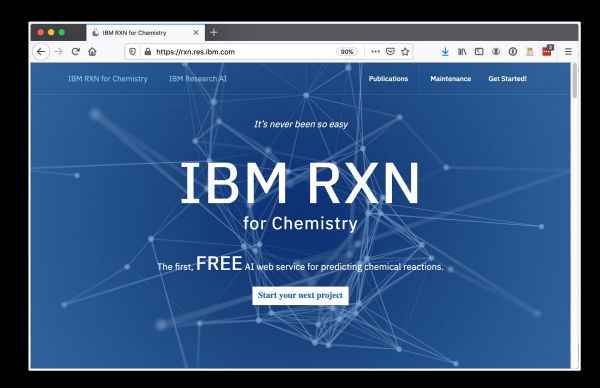


Summary





IBM RXN



Freely available on: **rxn.res.ibm.com**

Thank you for your attention!

If you have any questions:

E-mail: ava@zurich.ibm.com Twitter: @acvaucher Whova app

Acknowledgments:

Antonio CardinalePhilippe SchwallerAlessandro CastrogiovanniAleksandros SobczykJoppe GeluykensAlessandra ToniatoTeodoro LainoHeiko WolfMatteo ManicaFederico ZipoliVishnu H. NairFederico Zipoli

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