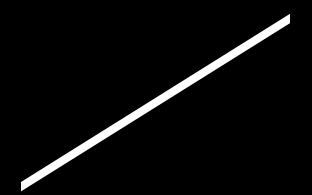
Learning from literatureextracted synthesis actions for organic synthesis





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

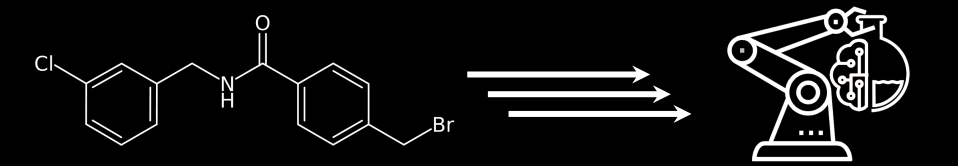




How can exploit this **data** to **accelerate discovery**?

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

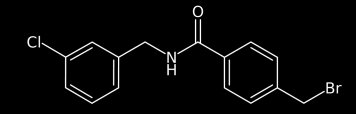
Data and chemical reactions



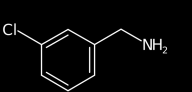
Target molecule

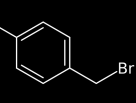
Synthesis execution

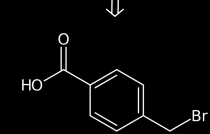
Step 1: retrosynthetic analysis



 \downarrow



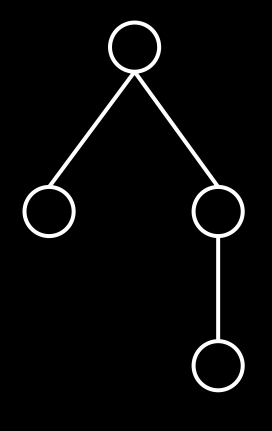




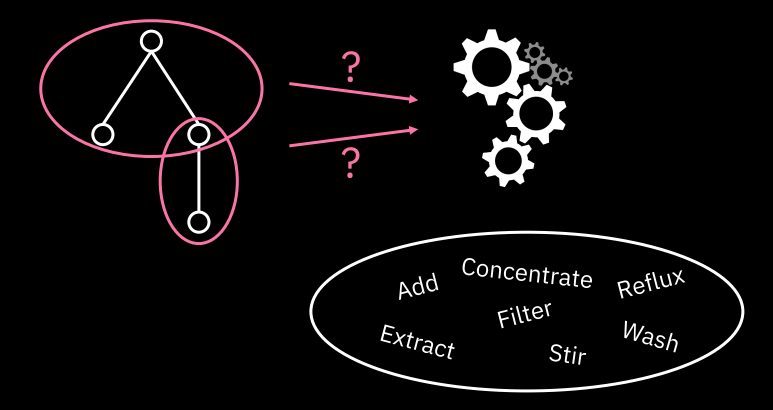
C

0

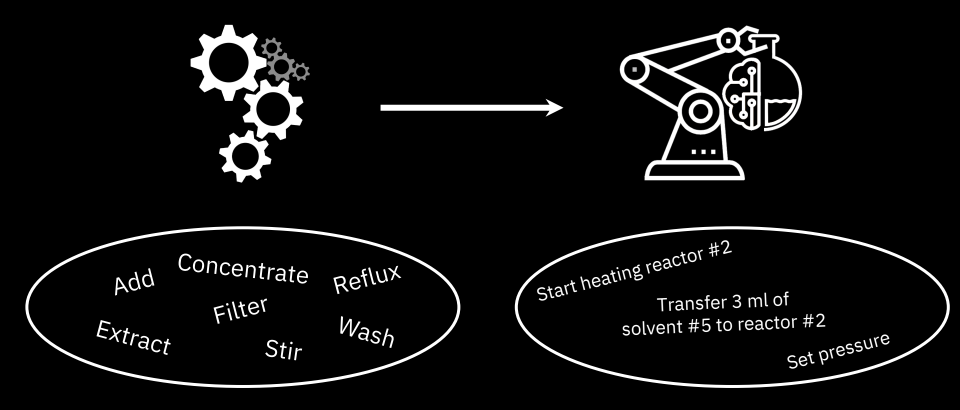
Retrosynthetic tree



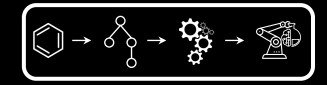
Step 2: experimental steps

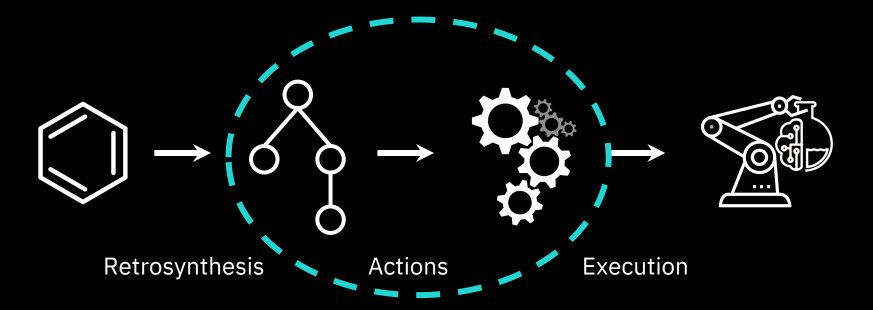


Step 3: Execution on robotic system

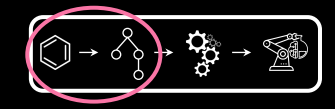


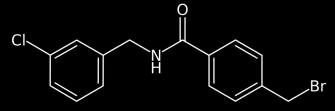
All together

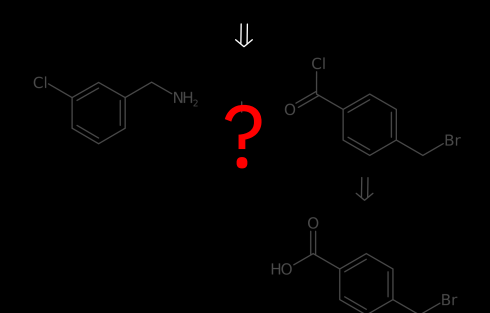


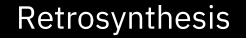


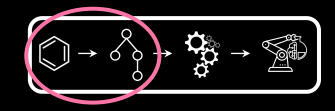
Retrosynthesis

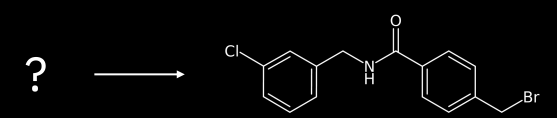


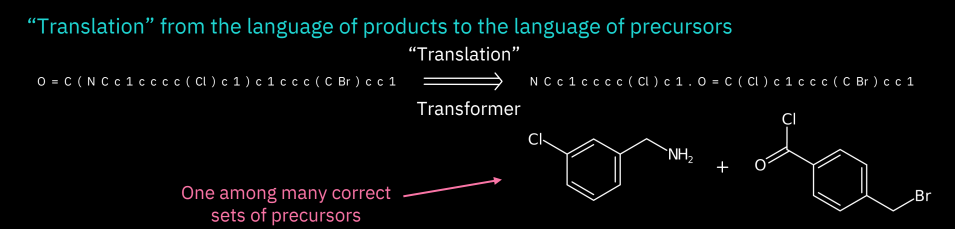






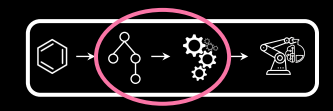


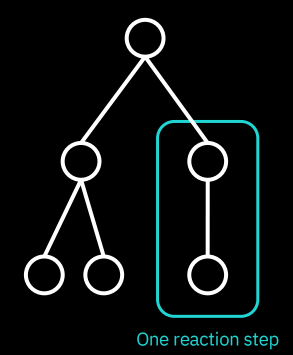


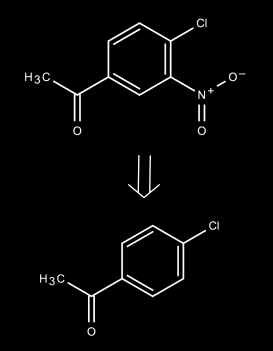


Schwaller, P.; Petraglia, R.; Zullo, V.; Nair, V. H.; Haeuselmann, R. A.; Pisoni, R.; Bekas, C.; Iuliano, A. & Laino, T., Chem. Sci., 2020, 11, 3316-3325.

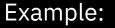
Synthesis actions

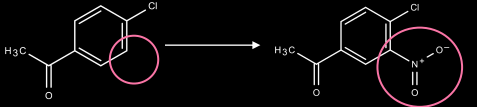


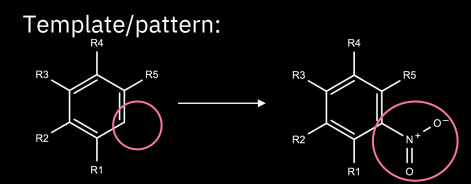




Synthesis actions



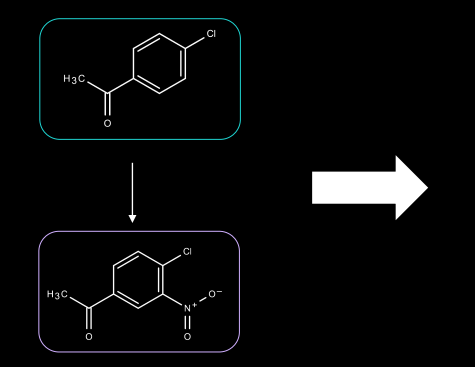




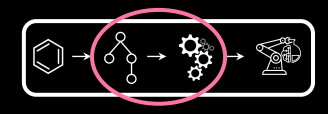


- -Same template but different synthesis actions!
- -Hard to predict
- –Ideally: ML model! "SMILES-to-actions"

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

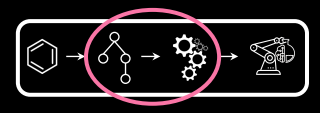
... Sodium borohydride (0.25 g, 6.6 mmol) was added portion wise over a period of 20 min and the reaction mixture was stirred for 3.5 hrs at 20-25° C ...





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	



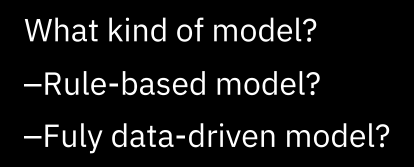
ction name	Variable name	Variable type
d	material	chemical
	dropwise	boolean
	temperature	string (optional)
	atmosphere	string (optional)
	duration	string (optional)
llectLayer	layer	string
ncentrate	(none)	
ystallize	solvent	chemical
gas	gas	string (optional)
	duration	string (optional)

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')



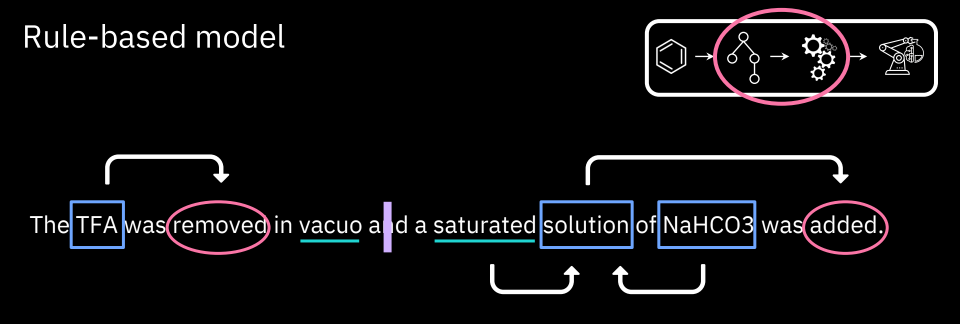
Models for Paragraph-to-actions



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:

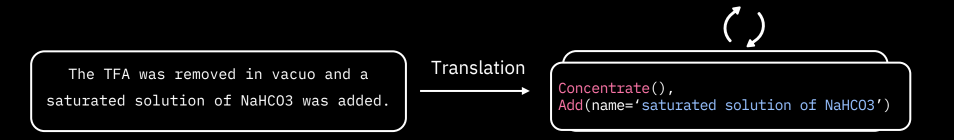
Rule-based training data for ML model



Generated actions for ~4M sentences

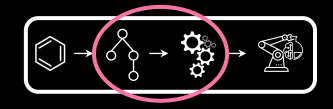
ML model





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.

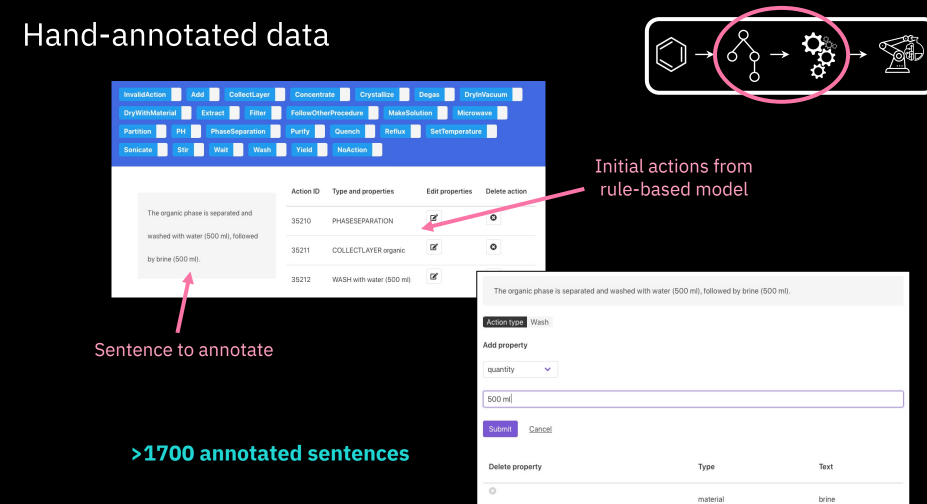
Rule-based model: ADD Diisopropylazodicarboxylate (0.05 ml, 0.302 mmol); STIR for 3 hours at room temperature. ML model: ADD Diisopropylazodicarboxylate (0.05 ml, 0.302 mmol); STIR for 3 hours at room temperature.

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

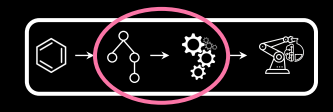
Rule-based model:CONCENTRATE; ADD water.ML model:CONCENTRATE; ADD water.Ground truth:CONCENTRATE; ADD water; PH with hydrochloric acid (1 M) to pH acidic.

Improving ML model:

- Training data size / quality
- Refine on human annotation

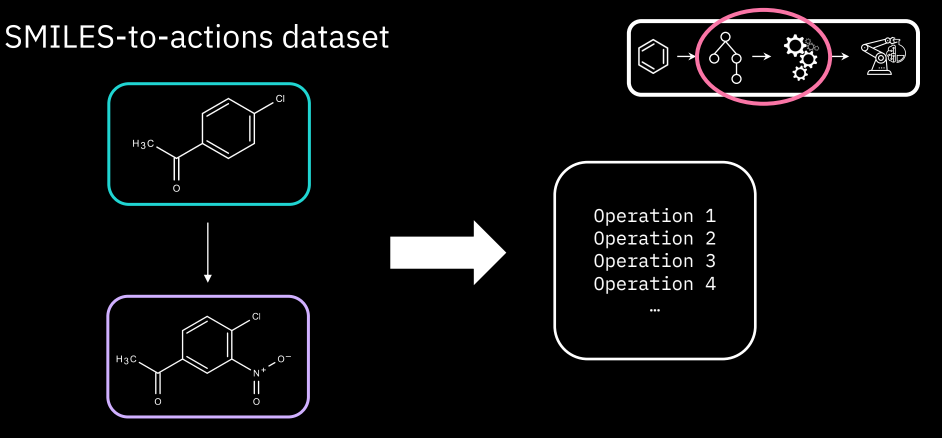


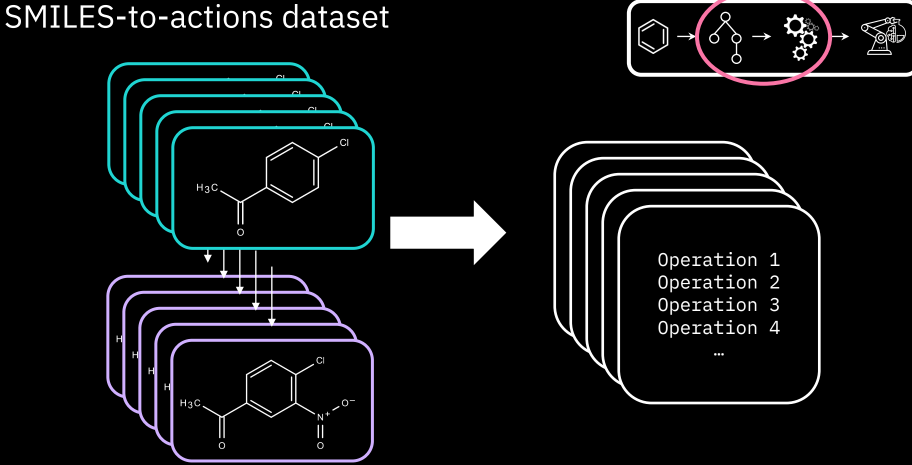
Results



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

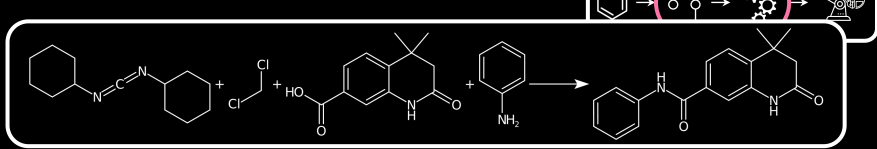
Vaucher, A. C.; Zipoli, F.; Geluykens, J.; Nair, V. H.; Schwaller, P.; Laino, T., *Nat. Commun.* **2020**, *11*, 3601.





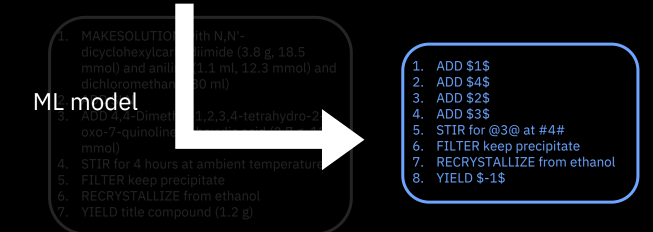
Vaucher, A. C.; Schwaller, P.; Geluykens, J.; Nair, V. H.; Iuliano, A.; Laino, T., *Nat. Commun.* **2021**, *21*, 2573. Alain Vaucher / IBM Research Europe / March 30, 2022

SMILES-to-actions



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

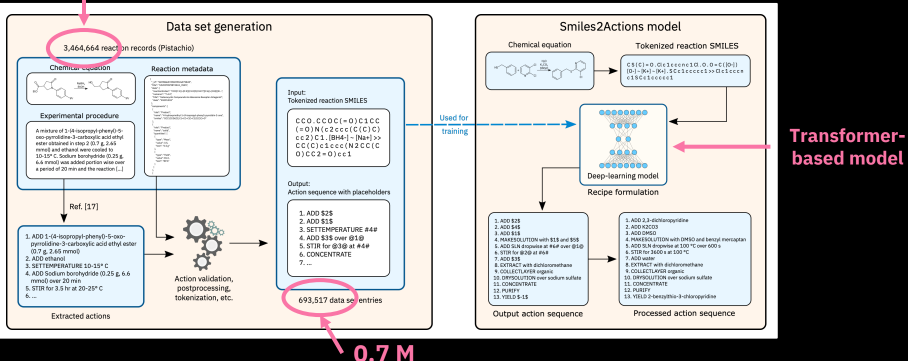
2.7 g (12.3 mmol) 4,4-Dimethyl-1,2,3,4tetrahydro-2-oxo-7-quinolinecarboxylic acid were added to a solution of 3.8 g (18.5 mmol) N,N'dicyclohexylcarbodiimide and 1.1 ml (12.3 mmol) aniline in 80 ml dichloromethane. The reaction mixture was stirred for 4 hours at ambient temperature and the precipitate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



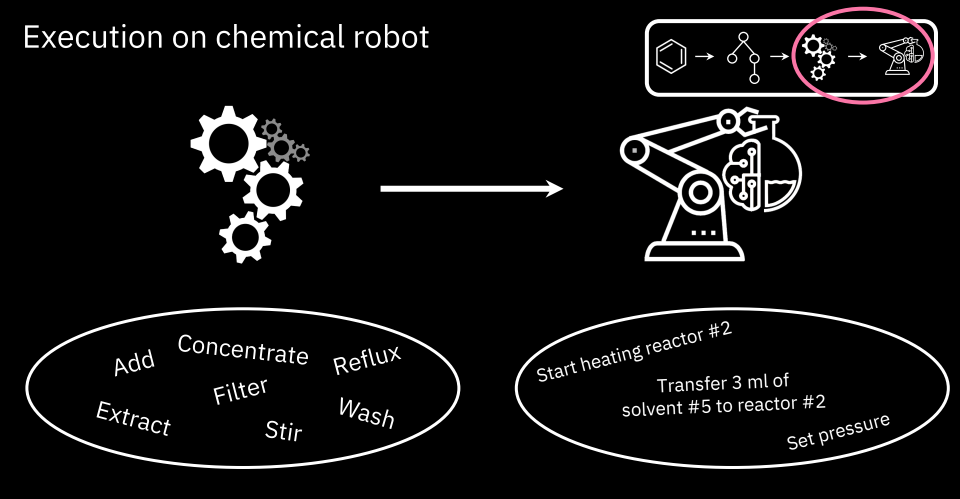
SMILES-to-actions

3.5 M

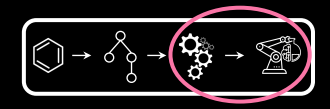




Vaucher, A. C.; Schwaller, P.; Geluykens, J.; Nair, V. H.; Iuliano, A.; Laino, T., *Nαt. Commun.* **2021**, *21*, 2573.



Execution on chemical robot



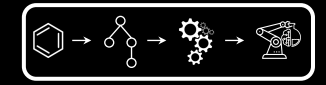
Cloud-based setup for autonomous synthesis

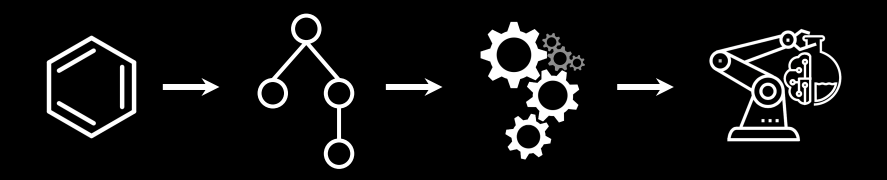


Hardware @ IBM Research Zurich

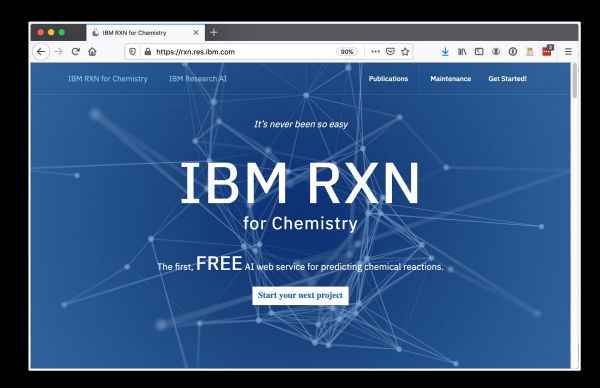


Summary





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

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

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