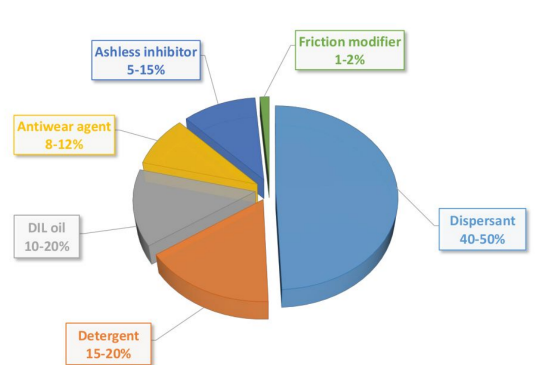


# Machine Learning for Molecular Design: a case study in dispersant design

Roi Naveiro SEIO - 2022

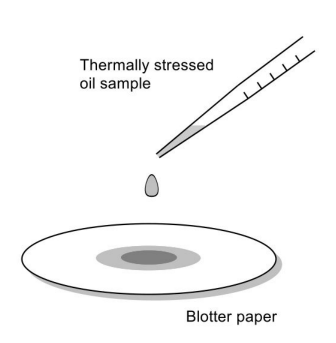
# **Dispersants in Lubricants**

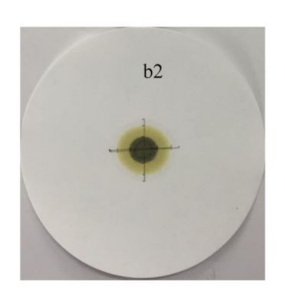


- Lubricants for combustion engines require formulated additive package (dispersants)
- Under harsh operating conditions of engines, soot is produced.
- Soot aggregation increases lubricant viscosity causing corrosion, deposit formation...
- Dispersants are molecules that adsorbs onto the surface of ultrafine carbon deposit precursors reducing their aggregation.

Goal: find molecules with high dispersancy efficacy

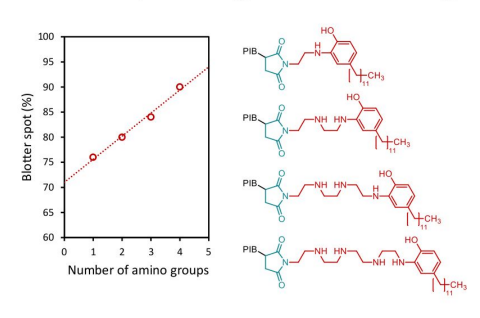
# **Measuring Dispersancy Efficacy - Blotter Spot**



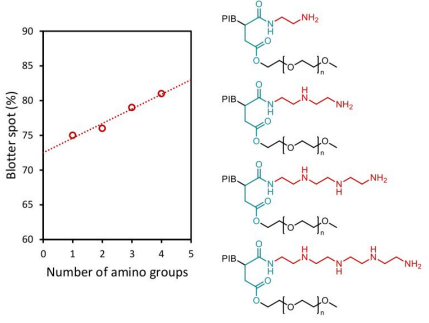


#### **Dispersancy Estimation - Limits of Chemist Intuition**

Within a family of substrate, predictable behaviors are appreciable.



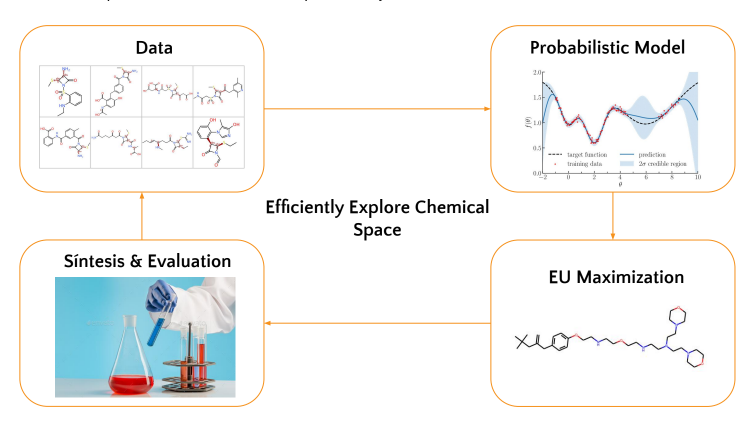
 However, the relationship between different families of substrates cannot be determined intuitively



Abdel Azim, A.-A. A. et al. *Int. J. Polym. Mater.* **2006**, *55*, 703 Abdel-Azim, A.-A. A. et al *Int. J. Polym. Mater.* **2007**, *57*, 114

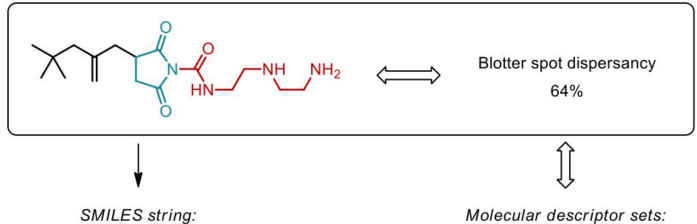
## Leverage data to find molecular structure with high blotter spot...

Solve black box optimization in chemical space (very limited number of evaluations!)



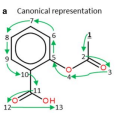
## **Probabilistic Model for Dispersancy - Data and Molecular Representation**

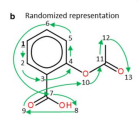
#### Dataset of 60 structures with associated Blotter Spot measure



O=C(C(CC(CC(C)(C)C)=C)CC1=O)N1C(NCCNCCN)=O

- Mordred package (425 descriptors)
- **SMILES embeddings** (769 descriptors)





#### **Probabilistic Model for Dispersancy - The Model**

- p >> N: sparsity inducing models
- Non linearity, interaction effects
- Bayesian Additive Regression Trees (BART): sum-of-trees model + regularization prior

$$y = \sum_{j=1}^m g(x; T_j, M_j) + \epsilon; \quad \epsilon \sim \mathcal{N}(0, \sigma^2)$$

Posterior inference through MCMC

$$p((T_1, M_1), (T_2, M_2), \dots, (T_m, M_m), \sigma | \mathcal{D})$$

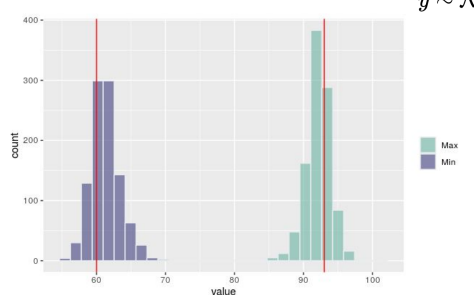
- Shallow trees capture varying (small) size interaction effects
- Natural way of performing variable selection (using variable importance measures)
- Better predictive performance than: linear regression with horseshoe prior, GP.

## **Probabilistic Model for Dispersancy - Prediction**

- ullet Given new structure with descriptors x, we need to sample from the predictive distribution  $\;p(y|x)\;$
- Sample

$$[T_j, M_j]_{j=1}^m, \sigma \sim p([T_j, M_j]_{j=1}^m, \sigma | \mathcal{D})$$

 $y \sim \mathcal{N}\left(\sum_{j=1}^{m} g(x; T_j, M_j), \sigma^2
ight)$ 

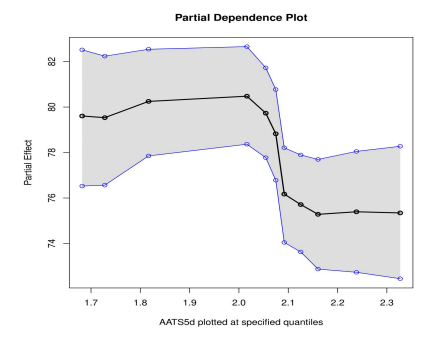


#### **EU Optimization**

- Idea: optimize expected utility to decide which structure to evaluate next
- Balance exploration vs exploitation
- ullet Expected improvement:  $\int \max{(y-y^*,0)\cdot p(y|x)dy}$
- ullet Probability of improvement:  $\int \mathbb{I}(y>y^*) \cdot p(y|x) dy$
- MC estimation
- How do we find structures that maximize a given expected utility?
- Difficult... rely on chemists!

## **EU Optimization - Interpretability**

- Chemist need to derive an actionable hypothesis from model output!
- ullet Provide partial dependence of each covariate in output:  $\mathbb{E}_{x_{-i}}\left[\sum_{j=1}^m g(x;T_j,M_j)
  ight]$

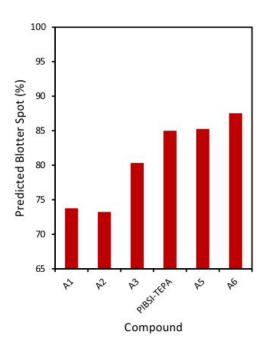


- But descriptors sometimes are difficult to interpret..
- In addition, some of the descriptors (neural embeddings) do not have interpretation!

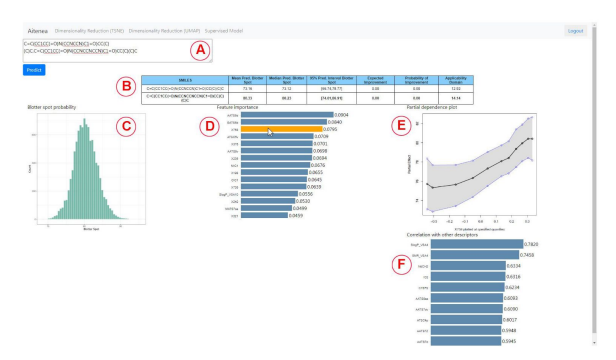
## **EU Optimization - Interpretability**

#### Validation and chemical interpretation

Density of amino groups in polar head



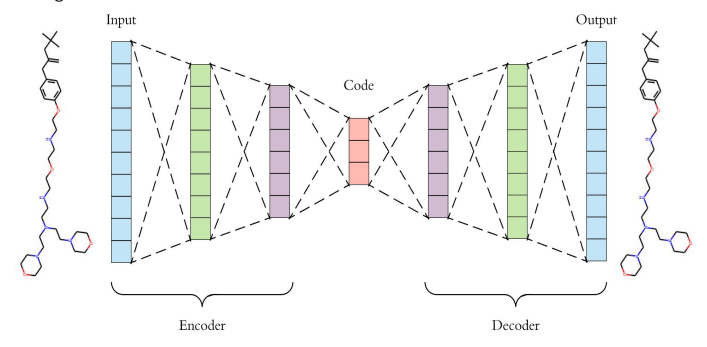
# **EU Optimization - Interpretability**



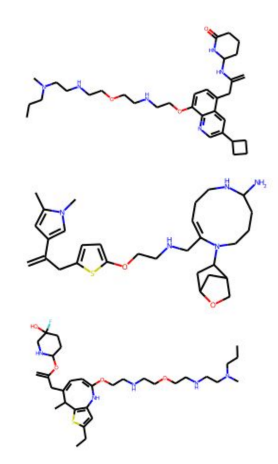
- Other trends discovered these way, allowed chemists propose molecules with good expected improvement
- Just one cycle of synthesis was enough for practical purposes...

#### Molecular Generation on a Nutshell

- Goal: generate molecules that maximize Expected Utility
- Several approaches depending mainly on algorithm and molecular representation
- Deep Learning based (VAEs)



# **Generation Issues**

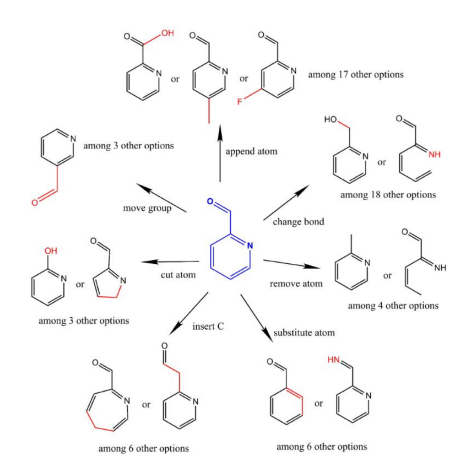


#### **Discussion**

- Statistical models can help accelerate molecular design
- Chemists need to interact with models. Interpretability is key (but very difficult)
- Removing humans from the process seems (almost) impossible. It would require automatic generation of new molecules
  - Multi-objective optimization
  - Small data regime
  - Structural constraints
  - Synthesizability
  - Uncertainty Quantification is key

#### Ongoing work

- Meta-heuristics for property optimization
- Genetic algorithms
- Iteratively mutate population of molecules (starting from a given one)



#### Acknowledgements

## CONICET













# Thanks!



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https://github.com/roinaveiro

# Why?







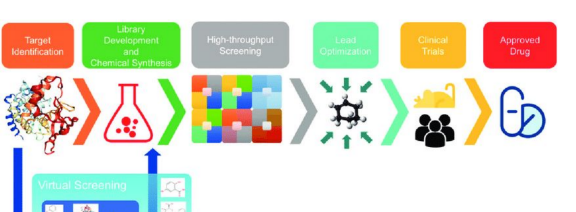


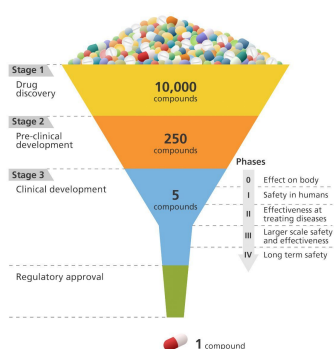
## **TOP 9 TAKEAWAYS**

Al investment in drug design and discovery increased significantly: "Drugs, Cancer, Molecular, Drug Discovery" received the greatest amount of private AI investment in 2020, with more than USD 13.8 billion, 4.5 times higher than 2019.

# The process of discovering new molecules

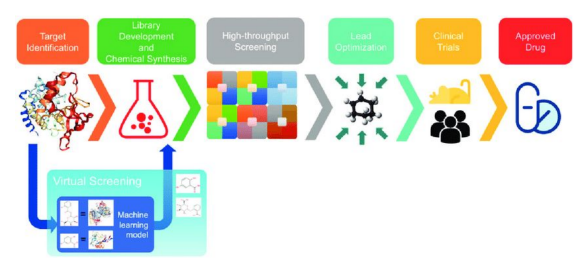
- Pharma: average time discovery market, 13 years
- Outside pharma: 25 years
- Crucial 1st step: generate pool of candidates
- Daunting task (e.g. 10<sup>23</sup> 10<sup>60</sup> drug-like molecules)





# The old way and the soon-to-be-old way

- Old way
  - Human experts propose, synthesize and test (in vitro)
- Soon-to-be-old way: high throughput virtual screening (HTVS)
  - Predict properties through computational chemistry...
  - ...leverage rapid **ML-based property predictions**



## De novo molecular design

- Just existing molecules are explored
- Much time lost evaluating bad leads
- Traverse chemical space more "effectively": reach **optimal molecules** with **less evaluations** than brute-force screening

"De novo molecular design is the process of automatically proposing novel chemical structures that optimally satisfy desired properties"



Combinatorial, black-box, stochastic, multi-objective optimization with black-box constraints

# Automatically proposing novel chemical structures

#### Two main ingredients

- Molecule representation
- Generative model

# Representing molecules

Molecules are **3D QM objects** with: nuclei with defined positions surrounded by electrons described by complex wave-functions

- Digital encoding that serves as input to model
- Uniqueness and invertibility
- Trade-off: information lost vs complexity
  - 3D coord. representation (symmetries?)
  - More compact 2D (graph) representation
- 1D, 2D and 3D

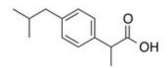


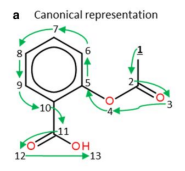
# **1D representations - SMILEs**

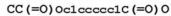
#### Simplified Molecular Input Line Entry System

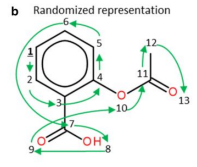
Molecule as graph (bond length and conformational info is lost)

- Graph traversal
- Sequence of ASCII characters
- Non-unique → Canonical SMILES
- One-Hot-Encoding
- Leverage NLP techniques
- SMILE-based methods struggle to generate valid molecules
- Valid = valency rules
- Learn spurious grammar rules



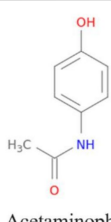




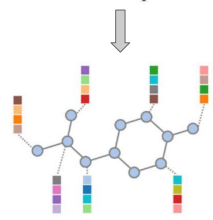


# 2D representations

- Nodes represent atoms
- Edges represent bonds
- Nodes/Edges have associated features (atom number, bond type, etc.)
- Capture connectivity!
- Symmetry invariant representation
- More difficult to generate than sequences
- Taylored algorithms that work with graphs (composing transformations on graphs, symmetries?)
- Graph Neural Nets!



#### Acetaminophen

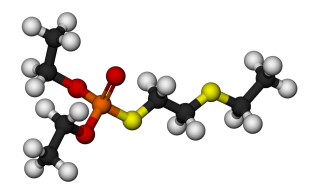


# 3D representations

• 3D point clouds

$$\mathcal{M} = \{x_i, r_i\}_{i=1}^p$$
 where  $x_i$  are features and  $r_i$  are coordinates.

- Minimal information lost (conformational preferences, bond lengths, etc.)
- Symmetries?
- Too many degrees of freedom
- Generation: sequentially choose pair of atoms, relative position, bond length and angles



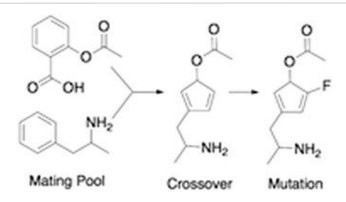
# How to generate molecules?

Myriad of different ways. A useful distinction:

- Gradient-free methods
- Gradient-based methods

#### **Gradient Free Methods**

- Graph-based genetic algorithms
  - Mutations and crossover on a pool of candidates
  - Elitist natural selection rule
- Yoshikawa et. al. propose using SMILES
  - Population of SMILES
  - Grammatical Evolution
- Many more...



## **Gradient Based Methods**

- Recurrent Neural Networks
- (Variational) Autoencoders
- Normalizing Flows
- Generative Adversarial Networks (GANs)

#### **Recurrent Neural Networks**

- Work on sequences (SMILES)
- Goal: given training sequences → learn to generate new sequences that resemble those of training.

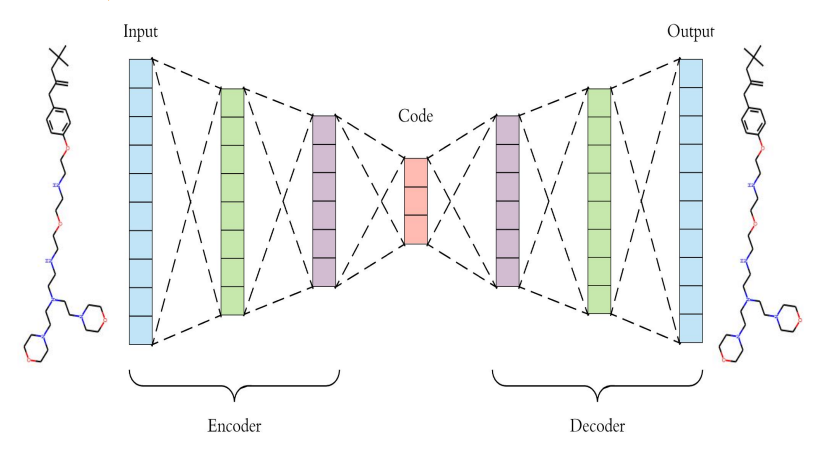
- ullet Sequence:  $S_{1:T} = (S_1, \dots, S_T)$  where  $S_i \in \mathcal{V}$
- Training: maximum likelihood, equiv to minimize loss function:

$$L^{MLE} = -\sum_{s \in \mathcal{T}} \sum_{t=2}^T \log \pi_{ heta}(s_t|S_{1:T-1})$$

- Generation: sequentially sample from multinomial dist.
- Thermal rescaling

$${\hat p}_i \propto \exp({p_i \over T})$$

# (Variational) Autoencoders



#### **Variational Autoencoders**

• Goal: learn probabilistic latent variable model for data generation

$$z \sim p(z) \ x \sim p_{ heta}(x|z)$$

ullet We want to maximize  $p(x)=\int p_{ heta}(x|z)p(z)dz$ ; instead maximize

$$\log p(\mathbf{x}) \ge \mathbb{E}_{z \sim q_{\phi}(z|\mathbf{x})} \left[ \log \frac{p_{\theta}(\mathbf{x}|\mathbf{z})p(\mathbf{z})}{q_{\phi}(\mathbf{z}|\mathbf{x})} \right]$$

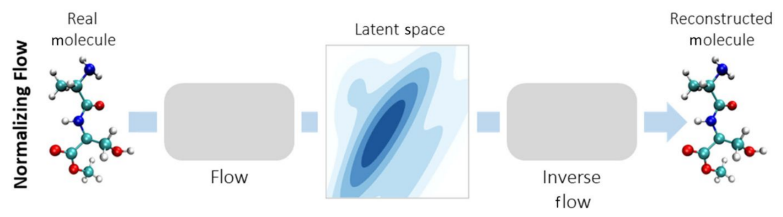
• RHS is equal to

$$\mathbb{E}_{z \sim q_{\phi}(z|x)}[\log p_{ heta(x|z)}] - D_{KL}[q_{\phi}(z|x), p(z)]$$

#### **Variational Autoencoders**

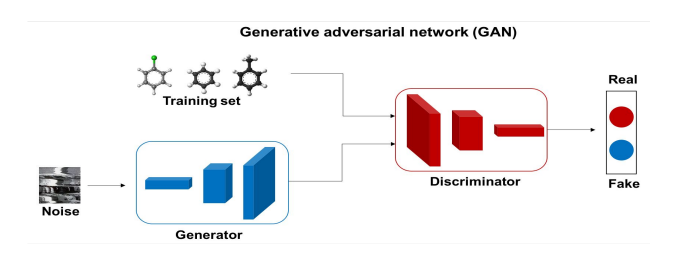
- Typically: p(z) independent standard normal dist. and  $q_{\phi}(z|x)$  factorized multivar. normal
- Mean and variance functions of encoder parameterized through CNN.
- Decoder normally RNN
- Training
  - Encode each training sample x into z
  - Decode z into x'
  - Minimize loss function
- Generation
  - Get point in latent space z
  - $\circ$  Decode z sampling  $\ x \sim p_{ heta}(x|z)$

# **Normalizing Flows**



- Learn series of parametric bijective transformations of probability distributions
- Allows (easy) calculation of exact likelihood.
- Deep NN with bijective layers

#### **Generative Adversarial Networks**



- Generator: generate molecule from Gaussian noise
- Discriminator: distinguish real from fake molecules
- Train to compete against each other

$$\min_{G} \max_{D} V(D,G) = \mathbb{E}_{\mathbf{x} \in p_{d}(\mathbf{x})} \Big[ \log D(\mathbf{x}) \Big] \\
+ \mathbb{E}_{\mathbf{z} \in p_{z}(\mathbf{z})} \Big[ \log \Big( 1 - D(G(\mathbf{z})) \Big) \Big]$$

#### Recall that...

"De novo molecular design is the process of automatically proposing novel chemical structures that optimally satisfy desired properties"



# Generate molecules that optimally satisfy desired properties

- Goal: learn valid molecules with desirable properties
- Infeasible to measure properties experimentally for every generated molecule...
- Infeasible to use computational chemistry to compute properties...
- Prediction: quantitative structure-activity relationship (QSAR)
- Done usually in separate datasets
- Many models depending on property, representation, etc.
  - Molecular Descriptors
  - SMILEs
  - Graphs

# Using properties to guide generation

- 1. Reinforcement Learning coupled with sequence generator
  - A time t, state is  $(s_0, \ldots, s_t)$
  - Action is next token  $a_t = s_{t+1}$
  - After taking action, a reward  $R_t$  is perceived
  - Goal, learn policy  $\pi_{ heta}(a|s)$

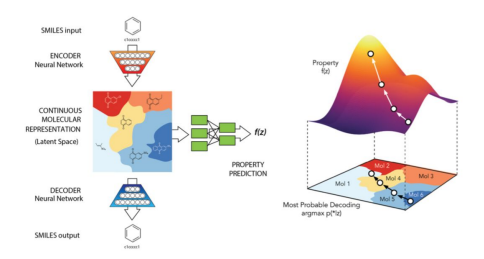
$$\max_{ heta} \mathbb{E}[\sum_{i=1}^T R_i | s_0, heta]$$

ullet The only non-zero reward is  $R_T$ which is equal to the property prediction

# Using properties to guide generation

#### 2. Optimization with VAE

- Learn map from latent space to property (e.g. through GP)
- Optimize that map (gradient ascent, bayesian optimization, etc.)



#### **Issues/Thoughts**

- Multi-objective optimization
  - Many properties to be optimized (depending even on different stakeholders!)
  - Drug discovery: high binding affinity to biological target, low toxicity, solubility, synthetically accessible, stability, economical costs!
  - Commonly: predict properties independently and combine predictions in loss function.
  - Also, hold properties constant implicitly through structural constraints.
  - Decision theory: multi-attribute utilities to incorporate different objectives for different stakeholders into the generative process

#### **Issues/Thoughts**

- Uncertainty quantification
  - Models rely on predictions to generate promising molecules
  - Accuracy of these models is key
  - o In small data regimes... models tend to be less accurate.
  - o Incorporate uncertainty quantification into generative process! (Bayesian inference)
  - Exploration vs exploitation (Bayesian optimization)
  - Bayesian decision theory

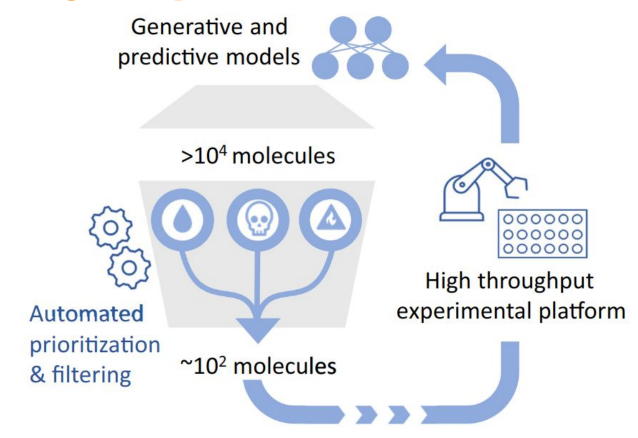
#### **Issues/Thoughts**

- Synthesizability
  - Generated molecules must be easy to synthesize
  - This concept is hard to define!
  - Methods to automatically evaluate synthesizability without human intervention
  - Rather than molecules, generate synthetic pathways (learn reactions)

#### Other relevant fields

- Graph based deep learning
- Geometric deep learning
- Combinatorial black-box optimization
- Heuristic search algorithms
- Reinforcement Learning

# The dream - Closing the loop



#### The reality?

- More likely: computer-aided molecular design
- Interpretability
  - Prediction is not enough, we need understanding (?).
  - Chemist need to derive an actionable hypothesis from model output.
  - If chemist sees, e.g. structural elements responsible for toxicity, she might have ideas on how to modify molecule to diminish toxicity
  - Interpretable representations: molecular descriptors...?
  - Interpretable methods to determine causality between structure presence and property (causal inference, counterfactual inference)

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### **Unconstrained generation**

- Goal: learn general distribution of molecules in chemical space
- Evaluated based on chemical validity, novelty, uniqueness