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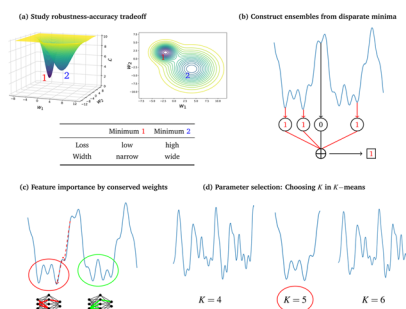


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Maximilian P. Niroomand, Luke Dicks, Edward O. Pyzer-Knapp* and David J. Wales*



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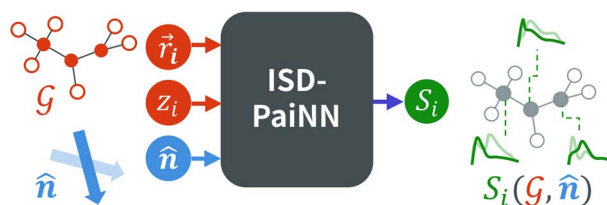
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Kiyoh Shibata* and Teruyasu Mizoguchi

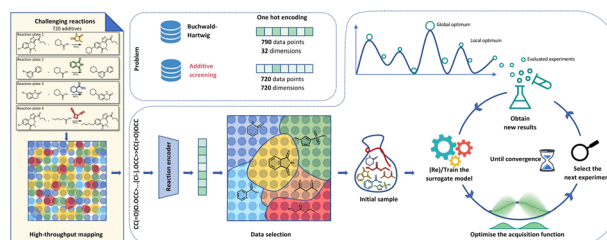


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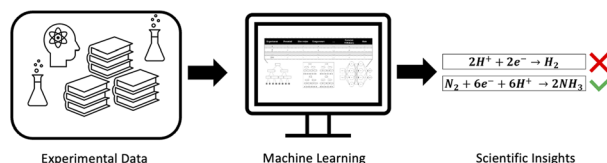
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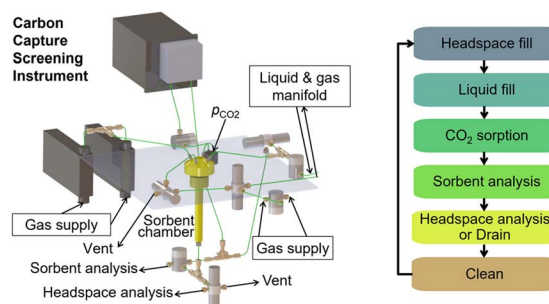
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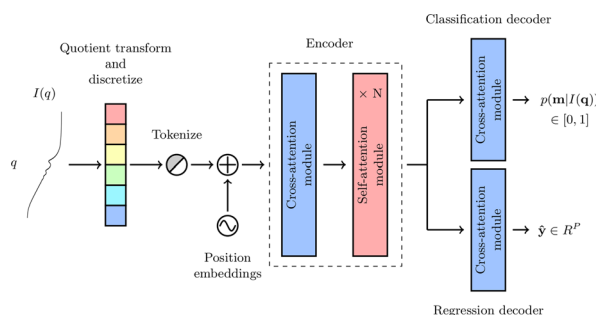
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Aleksei Krasnov,^{*} Shadrack J. Barnabas, Timo Boehme, Stephen K. Boyer and Lutz Weber^{*}

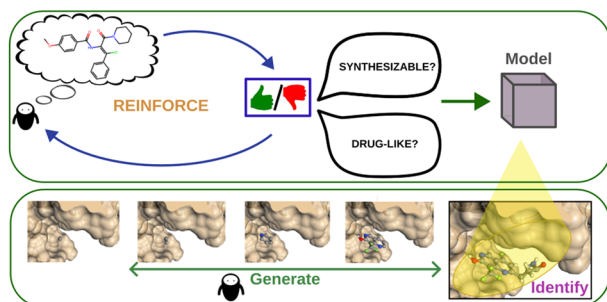
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Batuhan Yildirim, James Douth and Jacqueline M. Cole^{*}

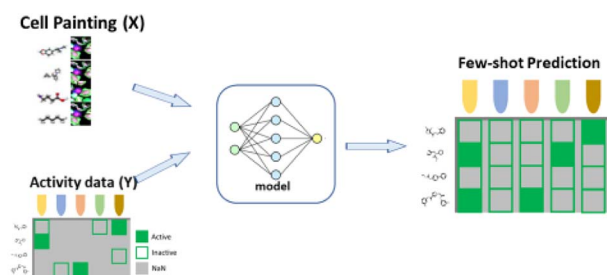
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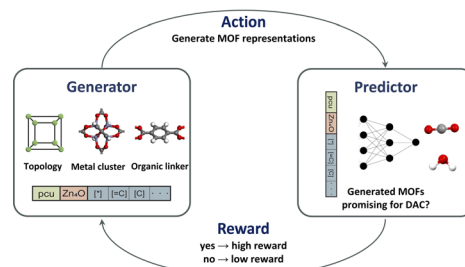


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Hyunsoo Park, Sauradeep Majumdar, Xiaoqi Zhang, Jihan Kim* and Berend Smit*

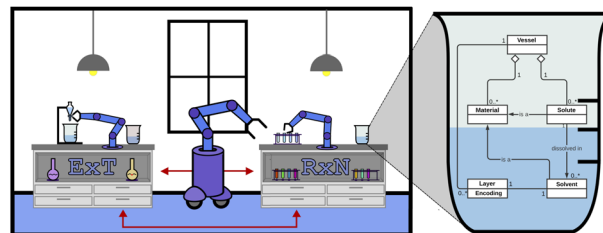
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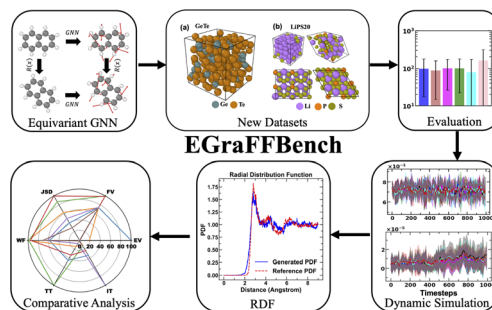
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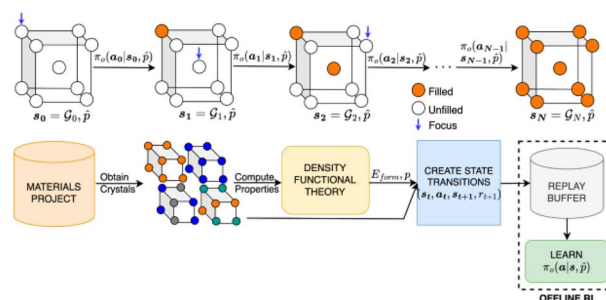
Vaibhav Bihani, Sajid Mannan, Utkarsh Pratiush, Tao Du, Zhimin Chen, Santiago Miret, Matthieu Micoulaut, Morten M. Smedskjaer, Sayan Ranu* and N. M. Anoop Krishnan*



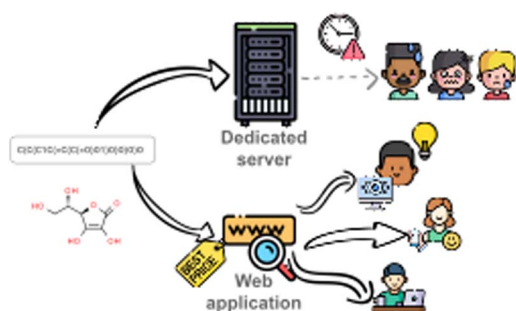
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Learning conditional policies for crystal design using offline reinforcement learning

Prashant Govindarajan,* Santiago Miret, Jarrod Rector-Brooks, Mariano Phielipp, Janarthanan Rajendran and Sarath Chandar



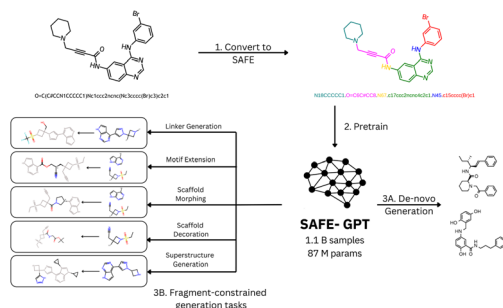
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Mayk Caldas Ramos and Andrew D. White*

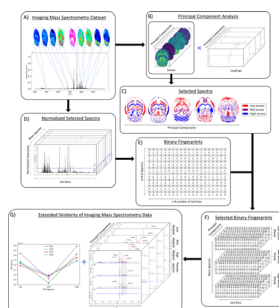
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Emmanuel Noutahi,* Cristian Gabellini, Michael Craig, Jonathan S. C. Lim and Prudencio Tossou

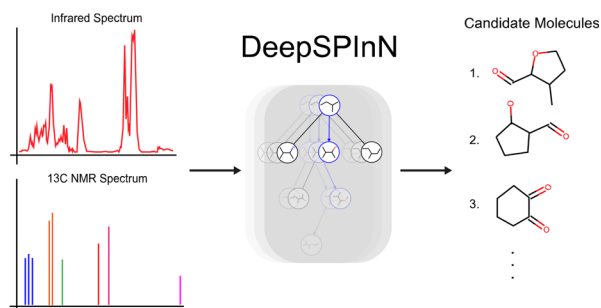
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Nicholas R. Ellin, Yingchan Guo, Ramón Alain Miranda-Quintana* and Boone M. Prentice*

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