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Subject: Re: Drug Likeness vs Drug Score  
Posted by [nbehrnd](#) on Mon, 01 Feb 2021 17:37:35 GMT  
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The manual's page

<http://www.openmolecules.org/properties/properties.html>

reads like a druglikeness is computed in first place, subsequently used to compute a drug score.

Lipinski's rule of five properties is one, but not the sole approach to define druglikeness. An implementation other than DataWarrior computing the later and others may be SwissADME (<https://doi.org/10.1038/srep42717>, open access).

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