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Subject: Re: Duplicate Molecules

Posted by [sbembenek18](#) on Wed, 04 Mar 2020 20:18:28 GMT

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I see what it is now. I have molecules where nitrogen might be considered a stereocenter. So, whereas the program that wrote the SDF considers that there are distinct stereoisomers, DW does not. I think this is consistent with the challenges of assessing this for these molecules.

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