
Subject: Re: How to customize Merge Equivalent Rows

Posted by [yuan](#) on Tue, 28 May 2024 01:40:52 GMT

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Hi Thomas,

Thank you for your reply. I downloaded the updated .jar file and tested it. Really fantastic!
I believe more users will benefit from it.

Yuan

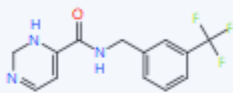
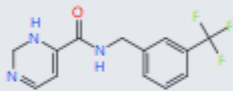
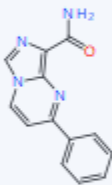
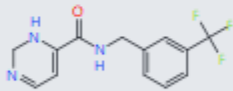
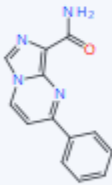
File Attachments

1) [Merge_results.png](#), downloaded 570 times

Test.dwar

File Edit Data Chemistry Database List Macro Help

Table

	ID	Structure of Structure	Structure	MS Assay	Species	T1/2(min)	Experimen Date
1	cmpd001		<chem>O=C(C1=CC=C(C=C1)C(F)(F)F)</chem>	MS	mouse	584.74	2024/1/24
2	cmpd001		<chem>O=C(C1=CC=C(C=C1)C(F)(F)F)</chem>	MS	dog	427.44	2024/1/24
3	cmpd002		<chem>NC(=O)c1nc2c(ncn12)c3ccccc3</chem>	MS	mouse	484.35	2024/1/24
4	cmpd001		<chem>O=C(C1=CC=C(C=C1)C(F)(F)F)</chem>	MS	monkey	153.73	2024/1/24
5	cmpd002		<chem>NC(=O)c1nc2c(ncn12)c3ccccc3</chem>	MS	dog	484.35	2024/2/24