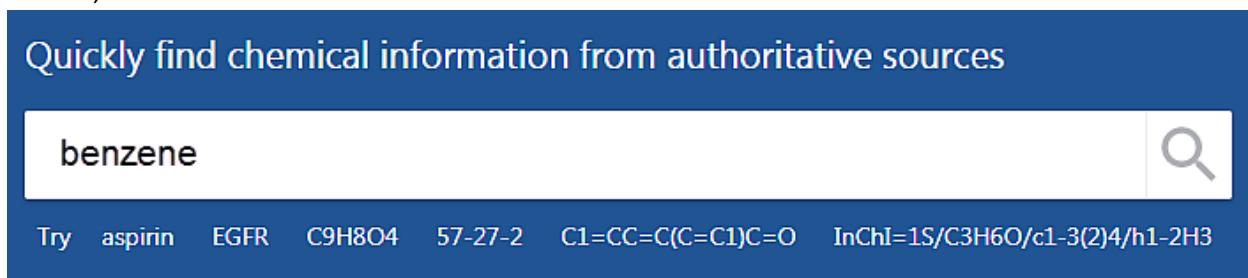


Accessing ChemIDplus Content from PubChem


ChemIDplus content is being migrated to PubChem. Follow the steps below to find the ChemIDPlus content in PubChem.

1. Start a search in the PubChem search by entering a chemical name, molecular formula, CAS RN, SMILES, or InChI identifier.



2. Select the “Best Match” compound

COMPOUND BEST MATCH



Benzene; Benzol; 71-43-2; Benzole; Cyclohexatriene; Pyrobenzole; Benzine; Phenyl Hydride; ...

Compound CID: 241
MF: C₆H₆ MW: 78.114g/mol
InChIKey: UHOVQNZJYSORNB-UHFFFAOYSA-N
IUPAC Name: benzene
Create Date: 2004-09-16

3. ChemIDplus content is distributed in various sections and fields in PubChem. Use the accordion style menu to expand/collapse the sections. Use “Ctrl F” shortcut key to find the field name.

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Title and Summary

1 Structures

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
1.2 3D Conformer

1.3 Crystal Structures

2 Names and Identifiers

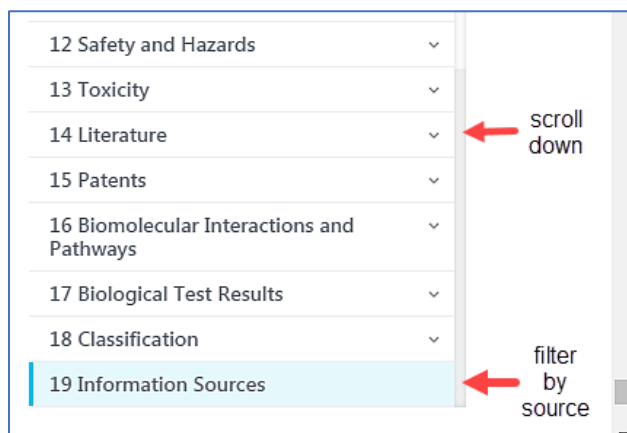
2.1 Computed Descriptors

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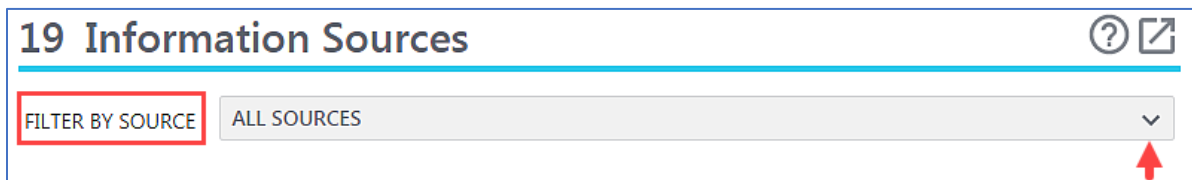


4. **IMPORTANT!** How to find **ChemIDplus-ONLY** content

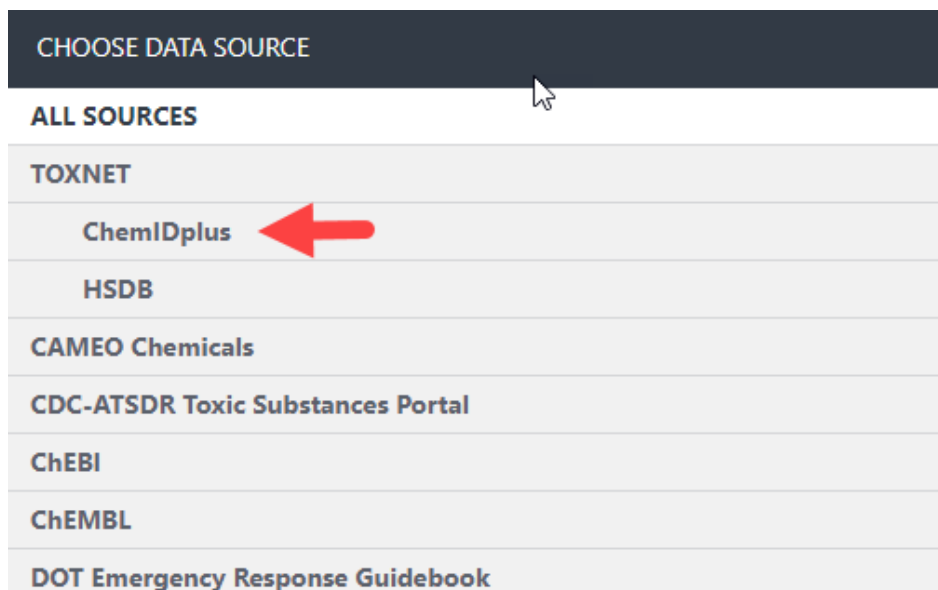
STEP 1: Scroll down to the **LAST** section “**Information Sources**”



STEP 2: Go to “Information Sources”, expand the “**FILTER BY SOURCE**” menu



STEP 3: Select **ChemIDplus** under the **TOXNET** sources.



Applying this filter will allow you to access ChemIDplus-only content in PubChem.