

Discovering new drugs is a long and expensive process: Chemical compounds that dupe screening tools make it even harder

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Modern drug discovery is an <u>expensive and complicated process</u>. Hundreds of scientists and at least a decade are often required to produce a single medicine. One of the most critical steps in this process is the first one—identifying new chemical compounds that could be developed into new medicines.

Researchers rely heavily on <u>bioassays</u> to identify potential drug candidates. These tests measure a compound's ability to act on a biological target of interest. Candidates that show up as a "hit" by interacting with a target of interest (such as fitting into a binding site on the target) move on to further study and development. Advances in technology called <u>high-throughput screening</u> have allowed researchers to run thousands of compounds through bioassays in a short time, significantly streamlining the process.

But some of these "hits" don't actually interact with the target as intended. And for the unwary researcher, this can lead down a rabbit hole of lost time and money.

I am a <u>medicinal chemist</u> who has been working in the <u>drug discovery</u> field for over 26 years, and one of the greatest challenges I have faced in my research is selecting good candidates from drug screening tests. One particular category of compounds, known as <u>pan-assay interference</u> <u>compounds</u>, or <u>PAINS</u>, is a common pitfall.

What are PAINS?

<u>Bioassays</u> involve placing a chemical compound together with the target of interest and measuring the strength of their interaction. Researchers assess interaction strength using a number of methods depending on how the bioassay is designed. A common assay design emits light when there is an interaction, where the intensity of the light depends upon the strength of interaction.



PAINS refer to compounds that often come up as <u>false positives</u> during the screening process. Because of certain characteristics of these molecules, they can interact with a target in nonspecific or unexpected ways. Some can even react chemically with the target. So while PAINS may come up as a hit in a screen, it doesn't necessarily mean they actually do what researchers hoped they'd do. <u>Common worst offenders</u> include compounds like quinones, catechols and rhodanines.

There are a number of ways that PAINS dupe bioassays.

Some PAINS have properties that cause them to emit light (or fluoresce) under certain conditions. Since many bioassays detect light as a signal for a hit, this can confuse the assay readout and result in a false positive.

Other PAINS can act as <u>redox cyclers</u> in bioassays—producing hydrogen peroxide that can block the target and be misread as a hit.

Similarly, some PAINS form <u>colloidal aggregates</u>—clumps of molecules that interfere with the target of interest by absorbing it or modifying the molecular structure. In rare cases, these clumps can even elicit a desired interaction with the target of interest because of their large size.

Trace impurities left over from manufacturing can also elicit a PAINS response.

To make things even more complicated, because PAINS react with targets much more strongly than most compounds that are true drug candidates, PAINS often appear as the most promising hits in screening.

What can be done about PAINS?

An estimated <u>5% to 12% of compounds</u> in the screening libraries academic institutions use for drug discovery consist of PAINS. Scientists



misled by a false positive can waste considerable time if they try to develop these compounds into usable drugs.

Since researchers became aware of the existence of PAINS, medicinal chemists have identified frequent offenders and actively remove these compounds from screening libraries. However, some compounds will always fall through the cracks. It is ultimately up to the researcher to identify and discard these PAINS when they show up as false positives.

There are a few things researchers can do to filter out PAINS. In some cases, visually inspecting compounds for structural similarities with other known PAINS can be enough. For other cases, <u>additional</u> <u>experiments</u> are necessary to eliminate false positives.

Testing for the presence of <u>hydrogen peroxide</u>, for example, can help identify redox cyclers. Likewise, adding detergents can help break up colloidal aggregates. And bioassays that do not use light detection to register hits can circumvent PAINS that emit light.

Even the most experienced medicinal chemist needs to be cognizant of the dangers of these false positives. Taking steps to ensure that these types of compounds don't make it to the next stage of drug discovery can avoid wasted time and effort and ultimately lead to a more efficient and cost-effective drug discovery process.

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