

Advances in the Application of Small-Molecule Probes in the Study of Cellular Signaling Pathways

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Abstract. Small-molecule probes are functional organic compounds of simple structure that can selectively recognize and report changes in specific biochemical molecules within the intracellular space. In recent years, with the rapid evolution of chemical biology, rationally designed small - molecule probes have become important tools for dissecting complex cellular signaling networks. For example, when it was the 1980s, Roger Y's laboratory. Tsiendeveloped calcium ion (Ca^{2+}) indicators such as BAPTA and Fura-2, which pioneered Ca^{2+} imaging technologies; Pan and colleagues emphasized in a recent review that progress over the past decade has been remarkable and that probes have been widely applied to the study of proteins, signaling pathways, and drug–target interactions. The present review comprehensively summarizes the definition and design fundamentals of small-molecule probes, paying attention to the structural design of the recognition moiety, reporter element, and linker area. It then studies major probe kinds, such as imaging probes, activity - based (active - site) probes, and bioorthogonal probes, together with representative cases. At last, it looks into the progress made in using small - molecule probes in kinase - related, Ca^{2+} and redox signaling pathways. By arranging representative studies and commenting on them, this review points up the value of small-molecule probes in bioimaging, proteomics, and biomarker discovery, and gives an outlook on emerging trends and challenges in probe technology.

Keywords: Small-molecule probes, Cellular signaling pathways, Fluorescent imaging, Activity-based probes

1. Introduction

As life - science research progresses towards more mechanistic and quantitative analysis, cellular signaling pathways are now seen as highly interconnected networks instead of linear cascades. For the dissection of such systems, tools are needed which can access endogenous components in their native contexts, note dynamic changes as time goes on, and resolve subcellular spatial structure. Traditional strategies—including antibody-based detection and genetically encoded reporters—offer strong specificity but often rely on fixation, endpoint measurements, or genetic manipulation, which may limit temporal resolution and practical applicability in certain cell types or tissues [1].

In view of this background, small - molecule probes have emerged as useful chemical tools to complement biological methods. In view of their small size, structural flexibility, and ability to pass

through membranes, these probes can be rapidly deployed, titrated by concentration, and used in different experimental systems for real - time monitoring of intracellular signaling events in live cells [2]. Besides reporting molecular dynamics, small molecules are able to selectively interfere with individual proteins or whole pathways, providing mechanistic understanding of complex phenotypes [3]. Unbroken progress in probe sensitivity and structural modification have further extended their applications in imaging, proteomics, and drug–target analysis [1].

In view of these trends, a systematic synthesis of probe design logic, classification, and pathway - focused applications is useful for guiding signaling research. This review thus sums up modular design principles, main functional categories, and typical applications in kinase, Ca^{2+} , and redox signaling, and then proceeds to discuss the advantages and future outlooks. The figure shows the whole conceptual framework. 1.

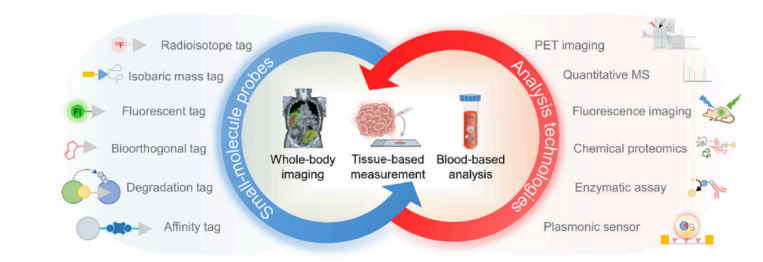


Figure 1. Overview of small-molecule probe modalities and their integration with analytical technologies for translational and clinical applications [1]

2. Concept and classification of small-molecule probes

2.1. Definition and design principles

This review describes small - molecule probes as organic compounds with a relatively low molecular weight (usually not exceeding 1000 daltons (Da)). Structurally, they are commonly described as comprising three basic elements: (1) a recognition moiety that selectively binds to a target molecule; (2) a reporter moiety (for example, a fluorescent dye or radioisotope) that generates an output signal; and (3) a linker region that connects recognition and reporter elements in a way that preserves both target engagement and signal generation [4].

In comparison to macromolecular probes such as antibodies or fluorescent proteins, small - molecule probes often have a more basic structure, lower cost, and are more easily synthesized and modified [4]. Because of their small size, they can facilitate cell entry and access binding sites that larger reagents may be sterically precluded from. Their synthetic tractability makes iterative optimization possible. A well - known case is the development of Ca^{2+} probes by Tsien's group. BAPTA was constructed by introducing aromatic rings into an ethylene glycol-bis(β -aminoethyl ether)-N, N, N', N'-tetraacetic acid (EGTA) scaffold to improve Ca^{2+} selectivity, while Fura-2 was produced by combining BAPTA with an aromatic fluorophore so that Ca^{2+} binding produces a detectable blue shift in the absorption peak [5]. This example suggests a general idea: probe behavior can be engineered by looking at recognition and reporting functions as modular building blocks and adjusting their connection through scaffold and linker design.

In excess of module selection, probe design must satisfy a number of practical criteria, especially for live - cell applications. A probe ought to be as non - interfering as reasonable, keep proper stability in physiological circumstances, and display high selectivity and sensitivity to its target. In respect of fluorescent probes, successful designs should stay stable under physiological conditions

with little photobleaching, bind their targets specifically, and give rise to a prominent optical response, ideally with low detection limits and wide dynamic ranges [4]. These criteria are not simply chemical. They judge whether readouts can be quantitatively analyzed in the face of biological complexity, cellular heterogeneity, and optical background.

A further good point about many small - molecule probes is that they're commonly used at micromolar or lower concentrations, helping to minimize perturbation of cell functions. This property can support applications not only in cultured cells. In some cases, it is also applicable to more complex biological models where keeping physiological states stable is important for meaningful interpretation. In practice, researchers can refine probe behavior by modifying the recognition moiety (to tune affinity/selectivity), selecting reporter elements (to tune emission wavelength, photostability, or modality), and optimizing linkers (to preserve binding while supporting signal transduction), thereby adapting probes to specific experimental needs.

2.2. Major types of small-molecule probes

According to design targets and experimental situations, small - molecule probes can largely be split into imaging probes and "omics - oriented" probes. The latter can be further divided into types like activity-based (active-site) probes and bioorthogonal probes. Despite the overlap of these categories in practice, the classification is useful as it links chemical mechanism to biological readout [6].

Imaging probes are designed so as to create directly observable signals—commonly fluorescence, but radioactive or other modalities too—upon binding to a target or after a certain chemical reaction. Their significance is in allowing real-time visualization of signaling molecules and activities in living organisms. Fluorescent imaging probes are able to either bind ions (e.g. Ca^{2+}) or experience chemical alterations according to the target which change optical properties. Examples of Ca^{2+} indicators, Fura-2 and Fluo-4, have fluorescence readouts that show the dynamic intracellular Ca^{2+} concentrations. Commonly employed probes such as 2', 7'-dichlorodihydrofluorescein diacetate (DCFH-DA) and Peroxyfluor-1 are used to find out the levels of reactive oxygen species. In imaging scenarios, probe design generally tries to increase signal - to - background ratios and allow for measurements over physiologically relevant concentration ranges and timescales.

In a different light, activity - based (active - site) probes are designed to covalently change a target class of enzymes by reacting with an active - site residue, thus reporting enzymatic activity rather than just protein quantity [7]. These kinds of probes include a reactive group which forms a covalent bond for the catalytic center of a specific enzyme family. A cited representative example in this review is the fluorophosphonate - biotin (FP - biotin) probe, used to label active sites of serine hydrolases. Once labeling has been carried out, affinity enrichment and mass spectrometry identification can make known the enzymes that were active (and so probe - reactive) under the tested conditions. The workflow enables chemical - proteomics analyses that can give information about activity states, count target classes, and assist in the discovery of new biological targets.

Bioorthogonal probes, commonly, contain functional groups (e.g., azides or alkynes) that make possible subsequent "click" reactions. After an initial probe - target engagement, a secondary labeling step may follow. In this step, a fluorophore or enrichment tag is added by way of bioorthogonal chemistry. A standard design idea is to put a photoactivatable (photoreactive) group and a bioorthogonal handle into a well - known small-molecule inhibitor matrix. In living cells, light can activate the cross - linking of the inhibitor to its target protein. Then, click chemistry can attach enrichment labels for pull - down and target identification. These design schemes extend the usefulness of probes from live - cell imaging and kinetics to downstream protein identification and mechanism - based analyses.

These kinds of probes are in a complementary situation. Imaging probes give maps of cellular signals with spatiotemporal features and are right for dynamic monitoring. By way of contrast, activity-based and bioorthogonal probes are very useful for identifying targets, validating drug mechanisms, and doing proteomics-level analyses. The availability of probe designs with diverse mechanisms enables researchers to pick strategies corresponding to their questions, balancing the needs of quantitative imaging against qualitative discovery goals.

3. Applications of small-molecule probes in cellular signaling pathways

3.1. Kinase-related signaling pathways

Protein kinases take up a central position in signal transduction. There is an encoding of hundreds of protein kinases in the human genome. These kinases are important members of major signaling cascades, like the MAPK and PI3K/Akt pathways. Kinases control essential biological processes like proliferation, differentiation, and metabolism via phosphorylation-dependent regulation. This review describes that small-molecule probes have two main functions for kinase signaling research: allowing for localization and dynamic monitoring, and enabling chemical control to investigate function.

Fluorescent probes can make imaging possible by binding to or labeling specific kinases. A representative approach is to couple a known kinase inhibitor to a fluorophore, making use of the inhibitor - kinase affinity to send a fluorescent signal to the target [8]. Zhang and colleagues attached inhibitors of the kinases Mps1 and PLK1 to a boron-dipyrromethene (BODIPY) fluorescent dye, making probes which could image these kinases in living cells. In mitosis, these probes are able to monitor spatiotemporal localization and activity patterns without the requirement of transgenic tags or antibody staining [9]. This approach may be of great use when genetic manipulation is difficult or when endogenous conditions are preferred for mechanistic understanding.

Related research has associated inhibitors or substrate - mimicking peptides of other kinases to fluorescent groups, which is useful for cell - cycle analysis and the study of aberrant signaling in cancer and other diseases. While the implementation details change according to various kinase families, the conceptual benefits remain stable. First off, in situ imaging gives direct evidence about where kinase - associated signals are in cells and how they change their distribution over time, often difficult to infer from bulk biochemical assays [10]. Second, since fluorescent probes can be used in the setting of living cells, they can support time - resolved analysis of pathway dynamics under perturbations such as drug treatments or signaling stimuli.

Other than just imaging, activity - based and bioorthogonal strategies are useful for kinome - scale ("kinase omics") studies [11]. As an illustration, kinases can be captured from cell lysates by means of probe constructs based on ATP analogs or inhibitor scaffolds. After a biotin tag is attached through click chemistry, affinity capture can enrich bound kinases for identification and profiling. Such strategies, usually called "Kinobeads" approaches, have had wide - spread applications in kinase inhibitor target discovery and the research of resistance mechanisms. The fundamental principle is that chemically engineered affinity reagents can convert binding interactions into detectable signals, facilitating systematic comparisons of kinase binding across compounds and conditions.

Besides, the ideas of bioorthogonal and photo - control have been applied for the exact manipulation of kinase pathways. This review brings forward a representative example. Chen and colleagues made use of the bacterial effector protein OspF, which shows high specificity for the dephosphorylation process of MAPKs like extracellular signal-regulated kinase (ERK). By

introducing photocleavable or chemically cleavable non-natural amino acids into OspF, the effector can be rendered "dormant" until triggered by light or a chemical stimulus, after which OspF regains activity and can suppress ERK signaling at defined times and subcellular locations (e.g., nucleus or cytoplasm) [12]. The chemical - trigger terminology has been flagged as being of uncertain nature; see Appendix. All things considered, this example demonstrates how chemically engineered control modules can be integrated with signaling effectors to get spatiotemporal regulation that conventional inhibitors alone can't manage.

In sum, small - molecule probes that are able to label, capture, or control kinases offer chemical toolkits for investigating kinase topology, dynamics, and function. These probes can support mechanistic analyses of kinase pathways and inform the design of targeted therapeutics by making possible live-cell imaging, proteome-wide target identification, and spatiotemporally resolved perturbation [9].

3.2. Calcium ion signaling pathways

Calcium (Ca^{2+}) ranks among the earliest identified signaling ions in cellular biology. Its clear spatiotemporal fluctuations form an adaptable signaling language that takes part in various physiological processes, including neuronal excitability, muscle contraction, and immune responses. The establishment of small-molecule Ca^{2+} probes, especially fluorescent indicators, is a fundamental element in chemical biology and bioimaging.

The source review notes that pioneering work by Tsien's laboratory in the late 1980s suggested that introducing "bisphosphonate" groups into a "styryl" framework to create a coordination motif could yield highly selective Ca^{2+} binding (terminology flagged; see mapping table and Appendix). Tsien made BAPTA in 1980 and used it as a basis to develop fluorescent Ca^{2+} indicators, starting a new period for observing intracellular Ca^{2+} dynamics. Subsequently, the group developed Fura-2 (described in the source as a membrane-permeant "diacetate" form; alternatives such as "AM ester" exist in English usage), which exhibits a shift of its absorption peak toward shorter wavelengths upon Ca^{2+} binding. Importantly, Fura - 2 permits quantitative Ca^{2+} imaging by getting ratios from dual excitation wavelengths, decreasing signal bias caused by disparities in probe concentration and non - uniform illumination. The source review has Figure 2 that shows the molecular structures of Ca^{2+} -free and Ca^{2+} -bound Fura - 2 and the corresponding absorption shift features [5].

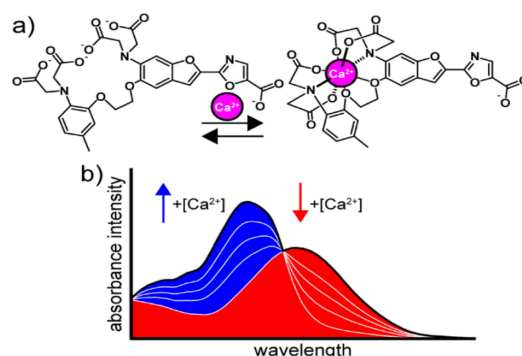


Figure 2. Structure and spectral changes of the Fura-2 calcium probe: a) Molecular structures of free Fura-2 and Ca^{2+} -bound Fura-2; b) Absorption spectral changes before and after Ca^{2+} binding (blue shift of the absorption peak after binding) [5]

Using similar design principles, Tsien's group further connected BAPTA-type chelators to fluorescein and rhodamine scaffolds, generating a series of single-wavelength, intensity-increasing Ca^{2+} probes such as Fluo-1/2/3 and Rhod-1/2. Among them, Fluo-3 and its derivative Fluo-4 are widely used for live-cell Ca^{2+} imaging as they provide higher quantum yield and better cellular loading efficiency than Fura-2. When Ca^{2+} binds to these probes, there are pronounced increases in emission intensity, allowing for the direct visualization of Ca^{2+} transients in neurons, muscle cells, and other excitable or signaling - active cell types.

The source review specifically indicates that new Ca^{2+} probes have been developed for specialized imaging conditions, such as two - photon and near - infrared modalities, to increase imaging depth and reduce the harm of photodamage in tissues. Taken together, these changes show how improvements in fluorophore engineering, probe delivery, and readout strategies can expand Ca^{2+} imaging from cultured cells to more challenging physiological contexts.

Generally, the ongoing improvements in fluorescent small-molecule Ca^{2+} probes have greatly boosted Ca^{2+} signaling research by making possible the quantitative imaging of Ca^{2+} dynamics in more and more complex physiological situations. The design principles set forth in this area, such as modular chelation chemistry, strategically chosen fluorophores, and ratiometric or intensity - based readouts, give transferable conceptual schemas for probe development in other signaling contexts.

3.3. Redox signaling pathways

Reactive oxygen species (ROS) and reactive nitrogen species (RNS)—including hydrogen peroxide (H_2O_2), nitric oxide (NO), and peroxynitrite (ONOO^-)—play dual roles in cellular biology: they can act as signaling molecules at controlled levels while also contributing to oxidative stress when dysregulated. Small-molecule probes give convenient methods to detect dynamic changes of these species in cells and so help with mechanistic investigations of redox signaling [1].

The review shows that Peroxyfluor-1 (PF1) is a frequently chosen probe for H_2O_2 . PF1 makes use of a boronate - ester group. H_2O_2 can selectively oxidize and cleave it, and this causes the release (or unmasking) of the fluorophore to generate fluorescence. These reaction - based designs can present high sensitivity and fast responses, making it possible to detect H_2O_2 increases taking place on biologically relevant timescales. Meanwhile, lots of redox probes undergo non - reversible chemical changes after reaction. This can put a limit on repeated measurements or make it complex to interpret steady - state levels against flux.

Reversible probes have also been developed to tackle more long - lasting and slowly changing redox alterations. The review gives Real Thiol as an illustration. As seen in Figure 3, this probe combines with glutathione (GSH) by a reversible Michael - addition reaction. The fluorescence signal can equate with the intracellular GSH concentration, making real - time quantitative measurement of GSH levels possible [13]. In contrast, traditional irreversible thiol probes usually give only qualitative or endpoint information since they modify thiols in a permanent manner and can't easily report dynamic equilibria.

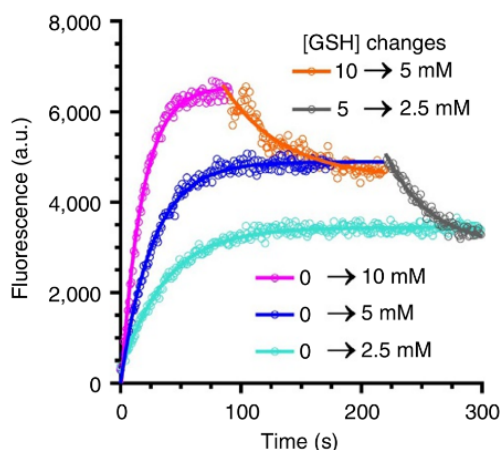


Figure 3. Reversible fluorescence response of the Real-Thiol probe to dynamic changes in glutathione (GSH) concentration [13]

In addition to H₂O₂ and GSH probes, the review notes that diverse probe frameworks have been developed for other ROS/RNS species, including NO probes based on acridine/hydrazone or thioether-oxidation mechanisms, and thiol probes based on oxidation of thioglycolic motifs, among others. Yet, since a lot of small - molecule redox probes change irreversibly once they react, their ability to capture subtle or highly transient signaling events may be restricted. Complementary approaches, say genetically encoded sensors (e.g. HyPer and such tools have been used to detect redox dynamics. But small - molecule probes remain important instruments because of their high sensitivity, simple delivery to cells, and compatibility with diverse detection methods [14].

4. Advantages of small-molecule probes

There are several advantages of using small molecule probes to study cellular signaling. Initially, they frequently show good cell permeability. Many probes are able to cross cell membranes and enable detection in living cells because their molecular weight is small [9]. As opposed to antibody staining or fluorescent protein tagging, small-molecule probes can usually be delivered without genetic engineering or physical transfection. This makes it possible to use them in primary cells, tissues, and other systems where genetic manipulation is a hurdle.

In addition, small - molecule probes have significant structural flexibility and can be modified chemically, at a fairly low synthetic cost. Researchers are able to introduce various affinity elements, fluorophores, or enrichment tags to customize probe functionality according to specific experimental requirements. The modularity makes possible rapid optimization cycles, which consist of adjusting target selectivity, signal modality, wavelength properties, or downstream compatibility with affinity capture and mass spectrometry workflows.

Apart from tuning target engagement, modular chemistry can also optimize physicochemical and signal - generation properties which strongly impact performance. For example, selection of longer-wavelength fluorophores can improve imaging under conditions where autofluorescence or limited light penetration would otherwise degrade signal quality, while incorporation of radioactive isotopes can enable whole-body readouts in appropriate modalities. Similarly, linker engineering is able to maintain the binding function of the recognition moiety and position the reporter for the greatest signal shift. As these parameters can be adjusted in an iterative way, small - molecule probes create a flexible platform for fitting tools to the constraints of specific signaling trials.

Thirdly, small - molecule probes typically bring about very little interference to the biological system under investigation. A good many probes produce measurable signals within the nanomolar - micromolar concentration range, and impacts on normal cellular function can be very slight [4]. In real - world practice, this low - dose operability is highly valuable in signaling studies. Pathway dynamics there can be sensitive to stoichiometric perturbations, and exogenous tools ought to be as unobtrusive as can be. Besides, ideal fluorescent probes are expected to present good biocompatibility and stable imaging performance in physiological conditions.

Lastly, small - molecule probes can be easily incorporated into existing analytical platforms, such as confocal microscopy, positron emission tomography (PET) imaging, and chemical proteomics. This multi - platform compatibility enlarges their application range and makes cross - validation between imaging - and proteomics - based readouts possible [13]. All in all, these advantages have made small - molecule probes necessary analytical tools for monitoring and disturbing signaling at molecular resolution in living - cell and in vivo circumstances.

5. Conclusions and outlook

To sum up, small - molecule probes have shown unique value as chemical tools for studying cellular signaling pathways. Starting with the design and classification of probes and going to applications in kinase, Ca^{2+} and redox pathways, the field has seen a consistent appearance of more sensitive and selective chemical probes. Fluorescent Ca^{2+} indicators and active - site labeling probes offer the capability to image signaling changes in real time at single - cell resolution. Meanwhile, reversible redox probes make dynamic and quantitative measurement of cellular redox states possible. These progressions not only heighten the basic understanding of mechanisms but also help with drug - target identification and the study of disease mechanisms, thus reinforcing the methodological basis for precision and translational research.

Going forward, it is expected that probe research will keep developing in the direction of multifunctionality and higher - level performance. One anticipated direction is closer integration with advanced imaging (including super-resolution fluorescence microscopy and multiphoton imaging) as well as improved synthetic methods such as late-stage functionalization and the design of near-infrared (NIR)-emitting dyes to enhance detection capabilities. Technologies that are just emerging, including artificial - intelligence - assisted molecular design, nanomaterial - enabled probe platforms, and microfluidic high - throughput screening, are also expected to make the process of probe development and optimization quicker [15]. As probe applications spread to more complex systems, focusing on biosafety, in vivo stability, and good readout performance stays important. It probably requires continuous chemical improvement and rational dose control.

In the presence of these opportunities, the practical application of probe technologies from cultured cells to more complex systems is still restricted by things such as biosafety and in - vivo stability. For actual applications, probes have to be stable enough to reach their targets under physiological circumstances and steer clear of long - term perturbations of cellular function. For reaction - based redox probes especially, irreversible chemical transformations, while making sensitive detection possible, can reduce the ability to capture subtle or highly transient signaling events [14]. To handle these constraints, it's probable that continuous chemical optimization and careful control of dosing and experimental conditions are needed, so that probe kinetics and signal dynamics are in harmony with the biological processes under measurement.

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