SAMDI Mass Spectrometry-Enabled High-Throughput Optimization of a Traceless Petasis Reaction

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Supporting Information

ABSTRACT: Development of the self-assembled monolayer/MALDI mass spectrometry (SAMDI) platform to enable a high-throughput optimization of a traceless Petasis reaction is described. More than 1800 unique reactions were conducted simultaneously on an array of self-assembled monolayers of alkanethiolates on gold to arrive at optimized conditions, which were then successfully transferred to the solution phase. The utility of this reaction was validated by the efficient synthesis of a variety of di- and trisubstituted allenes.

KEYWORDS: self-assembled monolayers, mass spectrometry, high-throughput experimentation, reaction optimization, allenes

The development of a synthetic methodology typically requires hundreds of reactions to be performed over a period of several months to identify effective combinations of reagents and to understand aspects of the structure−activity relationships. Several groups have reported progress with high-throughput approaches that significantly accelerate the development of reactions, yet this field is still in its early stages and requires further advances in methods that are efficient and applicable to a broad range of reaction types.1 Here we describe the use of SAMDI mass spectrometry2−6 to quickly perform and evaluate hundreds of variants of a traceless Petasis reaction,7 and we identify structure−activity relationships that in turn guided the development of the corresponding solution-phase reaction.

In 2012, we reported the development of a novel method for synthesizing allenes that involved the direct coupling of α-hydroxy aldehydes (i.e., 1) with alkynyl trifluoroborate salts (i.e., 2) in the presence of 2-nitrobenzenesulfonylhydrazide (NBSH, 3a) and La(OTf)3 (Figure 1, eq 1).7

While the reaction proceeded under mild reaction conditions to provide the desired allenes (i.e., 4) in good yield, attempts to expand the scope of the reaction to include substrates lacking α-hydroxyl substituents were unsuccessful. We viewed this shortcoming as an opportunity to explore the feasibility of conducting a high-throughput investigation of reaction parameters that would rapidly enable us to develop and optimize conditions to induce the desired transformation (Figure 1, eq 2). Ideally, the new reaction would proceed under mild reaction conditions, tolerate a range of aldehyde and ketone reaction partners (i.e., 5), and allow incorporation of a diverse set of substituents from the alkynyl trifluoroborate salt (i.e., 6), thereby affording access to a range of di- and trisubstituted allene products (i.e., 7) complementary to other methods.8

A few recent reports show exciting progress toward reaction screening.9−20 For example, the groups of MacMillan9 and Hartwig,10 have reported gas chromatography−mass spectrometry methods for analyzing hundreds of reactions. Porco and co-workers reported an analogous high-throughput platform based on liquid chromatography mass spectrometry,11 an approach subsequently improved upon in studies featuring ultra-performance liquid chromatography (UPLC)12 and Merck’s Multiple Injection in a Single Run (MISER) chromatography−mass spectrometry.13,14 Despite the limits

Figure 1. Traceless petasis reactions for the synthesis of allenes.
on throughput imposed by the chromatographic step, these methods have high generality in identifying products of interest. Other methods use DNA microarray, immunoassay, and label-assisted mass spectrometry formats but require assay-specific substrates that can be difficult to develop or that may limit the scope of substrates or conditions to be screened. Other tag-free approaches based on infrared thermographic techniques overcome these limitations but lack generality. Efficient and general methods should avoid synthetic tagging and slow processing steps (i.e., chromatography), be compatible with high-throughput formats and allow for the rapid evaluation of reaction products and yields.

In this work, we demonstrate a reaction optimization using the SAMDI array-based mass spectrometry approach. In the SAMDI method, chemical or biochemical reactions are performed on functional groups presented on a self-assembled monolayer. The monolayers are then analyzed with matrix-assisted laser desorption-ionization (MALDI-ToF) mass spectrometry to identify the masses of reaction products. Hence, SAMDI provides a label-free analytical tool for the characterization of each individual reaction. A further benefit of SAMDI is that reactions only need to be carried out on a picomolar scale, and the yields of these reactions can be quantitatively determined directly in a manner that obviates chromatographic purifications and synthetic labeling strategies. Here, we show how this method can be applied to determine optimal conditions that provide a high yield of the desired transformation in a single high-throughput experiment.

For the current study, we designed a high-throughput experiment that was based on 1800 unique combinations of 6 hydrazides, 15 alkynyl trifluoroborates, and 20 Lewis or Brønsted acid promoters (Figure 2A). These reactions were carried out on 6 arrays comprising steel plates having 384 gold spots each modified with a self-assembled monolayer of tri(ethylene glycol) terminated alkanethiolates. The carbonyl function required for the Petasis reaction was introduced on these gold islands by the direct Dess–Martin periodinane-mediated oxidation of the hydroxyl terminus (Figure 2B). We confirmed this transformation by analyzing the monolayers with SAMDI mass spectrometry, where the glycol-terminated monolayer showed peaks at m/z 693.6 and 709.6 (corresponding to the Na and K salts of the disulfide, respectively), and...
promoter (0.5 and 10 columns, and a unique Brønsted acid or Lewis acid of a 50 mM solution in acetonitrile) to 20 spots across 2 rows.

Representative example in Figure 2B, treatment of the NBSH were also analyzed by SAMDI mass spectrometry. For the internal standard. The resulting allene-functionalized plates known amount of an 11-amino acid peptide that served as an ethanol, followed by treatment with a THP matrix containing a and the plates were rinsed sequentially with water, acetone, and spot. The reactions were allowed to proceed for another hour, as.

where the oxidized monolayer showed a new peak at m/z 689.7 (Na salt of disulfide), as expected for generation of the aldehyde. The density of aldehyde functional groups was consistently more than 90% relative to the total alkanethiolates in the monolayer.

The next step in the workflow utilized automated liquid handling robots to deliver one unique arene-N-sulfonyl hydrazide 3a–f (0.5 μL of a 50 mM solution in acetonitrile) to 300 gold spots on each plate, leading to hydrazone formation after 1 h. Figure 2B shows a representative example using hydrazide 3a, where the SAMDI spectrum shows the generation of the expected hydrazone peak at m/z 1089.0, as well as an analogous peak at m/z 891.0 that shows the mixed disulfide wherein one end contains the hydrazone and the second terminus remained an unfunctionalized hydroxyl group. The six disparate hydrazone-functionalized plates were then rinsed with acetone and the robot was further used to sequentially deliver each alkynyl trifluoroborate 6a–o (0.5 μL of a 50 mM solution in acetonitrile) to 20 spots across 2 rows and 10 columns, and a unique Bronsted acid or Lewis acid promoter (0.5 μL of a 50 mM solution in acetonitrile) to each spot. The reactions were allowed to proceed for another hour, and the plates were rinsed sequentially with water, acetone, and ethanol, followed by treatment with a THP matrix containing a known amount of an 11-amino acid peptide that served as an internal standard. The resulting allene-functionalized plates were also analyzed by SAMDI mass spectrometry. For the representative example in Figure 2B, treatment of the NBSH (3a) derived hydrazone monolayer with trifluoroborate 6a and chlorotrimethylsilane led to a new peak at m/z 775.9 corresponding to the allene product. Automated software was used to extract the intensities of the expected allene product of each reaction as well as of the internal standard, and their ratio was computed and visualized through conditional formatting in a heat map for ease of analysis (Figure 3, see the Supporting Information for complete experimental details).

An analysis of the heat map shown in Figure 3 reveals that hydrazide 3a (NBSH) proved to be the most efficient for forming allenes on the self-assembled monolayer, although the sterically congested, electron–rich hydrazides 3e and 3f were also reasonably effective. The phenylacetylene trifluoroborate 6a generally provided the highest yields of the corresponding allene across the widest range of conditions, followed by the saturated and unsaturated cyclohexane derivatives 6i and 6j, and other aliphatic alkynyl trifluoroborates 6h, and 6k–l. Other aryl substituted alkynyl trifluoroborates 6f showed attenuated reactivity, though they were effective in the presence of HCl as the promoter. Based on the consistently high-yielding surface reaction between alkynyl trifluoroborate 6a and hydrazide 3a, we identified the six most efficient activators in the interfacial traceless Petasis reaction to be HCl, La(OAc)3, Sc(OTf)3, BF3·OEt2, TMSOTf, and TMSCl. We next evaluated these reagents in solution phase reactions using aliphatic aldehyde 5a (Table 1).

We found a reasonable correlation between the reactivities on the surface and in solution, but also noted several differences. For example, the Lewis acids Sc(OTf)3 and La(OAc)3 gave low yields of the allene for the interfacial reactions (Table 1, entries 2 and 4), unlike HCl and BF3·OEt2, each of which showed good reactivity on both the surface and the solution phase. TMSCl, in contrast, (Table 1, entry 6) gave better yields in solution than did several activators that performed better in the surface reaction. These changes are similar to those observed when solvent effects are examined and are not unexpected, since solvation at the interfacial region is distinct from that in the bulk and presents a unique reaction environment.
Ultimately, we found that BF$_3$·OEt$_2$ was the most effective activator for the solution reaction, providing the product in 58% yield using 1.5 equiv of alkyne 6a and 1.0 equiv of NBSH (3a). A related reaction involving the addition of lithiated acetylides into preformed N-aziridinylimines has been reported previously by Kim and co-workers, although only 12 examples were given and the reaction required 3.0 equiv of the alkyne reagent to generate good yields.$^{21}$ We therefore chose to further explore the efficiency of the BF$_3$·OEt$_2$-promoted reaction in solution (Table 2) to devise a more general and widely applicable method for allen e synthesis. The initial “hit” of 58% yield for the synthesis of allene 7a was readily increased to 69% yield when the amount of NBSH (3a) used was increased from 1.0 equiv to 1.5 equiv, and the temperature was lowered from room temperature to $-10$ °C. Attempts to decrease the amount of alkyne 6a or BF$_3$·OEt$_2$ used in the reaction led to diminished yields of the allene 7a, while increasing the amount of 6a beyond 1.5 equiv had no effect. Thus, the high-throughput investigation enabled by SAMDI, conducted on significantly miniaturized scales, allowed for the rapid discovery of effective reaction conditions, thereby significantly reducing the necessary amount of solution-phase optimization to only a few focused reactions.

Under the newly devised conditions, a number of allenes could be accessed from a variety of cyclic and acyclic aliphatic aldehydes and ketones (Table 2).

A range of linear, branched and cyclic aldehydes delivered disubstituted allenes in good to excellent yield upon reaction with phenylacetylene trifluoroborate 6a (Table 2, 7a–7f). Benzyl ether, alkenyl and alkyne functionalized aldehydes proved to be tolerant of the reaction conditions (Table 2, 7g–7l), with the formation of allene 7h demonstrating that terminal alkynes do not participate in the reaction (Table 2, 7h). For aldehydes possessing $\alpha$-stereocenters, the reaction proved to be poorly diastereoselective (Table 2, 7i and 7j). The alkylnyl trifluoroborate partner could also be readily changed, allowing access to a diverse set of aryl and alkyl substituted products (Table 2, 7k–7u). The oxygenated derivatives 7t and 7u were achieved in reasonable yields when the reactions were conducted at lower concentrations. The parent alkylnyl trifluoroborates for these substrates showed reduced solubility in acetonitrile, and an increased propensity toward unproductive protodeboronation under the standard reaction conditions.

Finally, we demonstrated that the traceless allene synthesis is also tolerant of both cyclic and acyclic ketones as reaction partners, allowing the generation of trisubstituted allenes in modest to high yields (Table 2, 7v–7x). The relatively mild reaction conditions, low reaction temperature and operational simplicity of this allene synthesis are particularly noteworthy, especially in the context of trisubstituted allenes, as many of the previously established methods require relatively harsh conditions and elevated temperatures.$^{25-24}$

In summary, we have demonstrated the application of SAMDI mass spectrometry for the high-throughput development and optimization of organic reaction parameters. By using the automated ability of the SAMDI mass spectrometry platform to rapidly conduct and analyze sequential reactions at miniaturized scale, we were able to prepare and analyze 1800 unique variations of a traceless Petasis reaction in order to provide initial reaction conditions and early insight into substrate scope. Translation of the most efficient surface-based conditions to the solution phase rapidly established an optimal set of reaction conditions to allow for preparative scale
A.B.D. thanks the ARCS Foundation, the University of Nebraska, and the National Science Foundation (CHE1361173 to R.J.T.). A.B.D. thanks the ARCS Foundation, the University of Nebraska, and the National Science Foundation (51). Allenes by a Traceless Petasis Reaction. Herein also set the stage for the further development and refinement of SAMDI mass spectrometry-based methods for future application to other organic reactions.

**REFERENCES**