High-speed Ball Milling Enables an Ultra-fast Wittig Olefination Under Ambient and Solvent-free Conditions Without Ylide Pre-formation.

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Abstract:

30 Seconds to success! – The Wittig reaction, a fundamental and extensively utilized reaction in organic chemistry, enables the efficient conversion of carbonyl compounds to olefins using phosphonium salts. Traditionally, meticulous reaction setup, including the pre-formation of a reactive ylide species *via* deprotonation of a phosphonium salt, is crucial for achieving high-yielding reactions under classical solution-based conditions. In this report, we present an unprecedented protocol for an ultra-fast mechanochemically driven Wittig reaction under solvent-free and ambient conditions, eliminating the need for tedious ylide pre-formation under strict air and moisture exclusion. A range of aldehydes and ketones were reacted with diverse phosphonium salts under high-speed ball milling conditions, giving access to the respective olefins in only 30 seconds.

Introduction:

Seventy years ago, Georg Wittig reported a ground-breaking reaction that revolutionized synthetic organic chemistry: the Wittig olefination.¹⁻³ This reaction, which converts carbonyl compounds into their unsaturated analogues *via* a [2+2] pericyclic mechanism, remains one of the most fundamental and extensively utilized transformations for the synthesis of functional alkenes.³⁻⁶ Despite numerous adaptations, the classical Wittig reaction continues to be crucial in modern synthetic chemistry, with applications ranging from polymer chemistry⁷⁻⁹ and material sciences^{10,11} to drug and natural product synthesis.¹²⁻¹⁷ Even on an industrial scale, the Wittig reaction is frequently pivotal; a prominent example is BASF's ton-scale synthesis of vitamin A and its derivatives.^{5,18-23}

In light of growing environmental concerns, there is an urgent need for more sustainable and greener modifications of this widely used reaction.²³ Recent efforts have focused on developing catalytic versions of the Wittig reaction, using strategies to regenerate phosphine *via* chemical or electrochemical reduction of the resulting phosphine oxide.²⁴⁻²⁸ Although these methods can recycle by-products, they often generate wasteful by-products themselves, such as when using chemical reductants like phenyl silane (PhSiH₃).²⁴ Additionally, attempts to use water as a reaction medium have been made,²⁹⁻³⁴ but the energy-intensive process of treating aqueous chemical waste often mitigates any environmental and cost benefits.^{35,36} Thus, the development of more sustainable and environmentally benign olefination processes remains a significant challenge.

Mechanochemistry has emerged as a transformative approach in this regard significantly advancing greener practices in academia and industry.³⁷⁻⁴⁰ This method uses mechanical force and friction to drive chemical reactions, often resulting in rapid reaction kinetics and either eliminating the need for solvents entirely or minimizing their use to a bare minimum—a strategy known as liquid-assisted grinding

(LAG).⁴¹⁻⁴³ Mechanochemistry frequently enables reactions that are highly air and/or moisture sensitive in solution to proceed smoothly under ambient conditions.⁴⁴⁻⁴⁶ These features make mechanochemistry an attractive and suitable strategy for modern synthetic chemistry.⁴⁷⁻⁵⁰

Despite its potential, solvent-free or LAG Wittig-type reactions are relatively underexplored in the literature (Figure 1). Most reports focus on stabilized phosphonium ylides in the Horner-Wadsworth-Emmons (HWE) modification of the Wittig reaction⁵¹⁻⁵⁶ or semi-stabilized ylides from benzyl phosphonium salts.⁵⁷⁻⁶⁰ While these reactions are conducted under neat conditions, some protocols still require external heating, such as conventional heating *via* an oil bath or microwave irradiation, to provide the necessary energy.^{51,53} Reports relying solely on mechanical force to drive the reaction are scarce.^{52,54,57-59,61}

In 2018, Mack explored the diastereoselectivity of the Wittig reaction on benzaldehyde under LAG using benzyltriphenylphosphonium bromide.⁵⁷ And very recently Friščić reported a hexameric supramolecular cage assembly constructed from (dibromomethyl)triphenylphosphonium bromide units.⁶¹ This assembly can encapsulate small carbonyl compounds and, upon mechanical impact, yield the respective 1,1-dibromoolefins *via* a base-activated host (PPh₃CHBr₂) - guest (aldehyde/ketone) directed Wittig olefination.

In 2002, Balema and Pecharsky explored the mechanochemical generation of phosphonium ylides under a helium atmosphere using K_2CO_3 as a base.⁵⁹ However, their study was limited, focusing on only three phosphonium salts reacted with three aromatic aldehydes. This investigation presented the exclusive example to date of a solvent-free methylenation reaction using a trimethylphenylphosphonium salt. Notably, employing 2-naphthyl aldehyde as a substrate, they obtained the respective vinyl derivative in a 73% yield after an extensive milling time of 20 hours.



Figure 1. Typical reaction conditions for the Wittig reaction of 4-phenyl benzyldehyde with PPh₃MeBr in solution (top, left), previous work on mechanochemical Wittig reactions by Balema, Mack, and Friscic (top, right), and our herein presented work with its advantages compared to reactions in solution (bottom, left) and selected examples thereof (bottom, right).

These findings led us to hypothesize that a Wittig reaction using non-stabilized phosphonium ylides under ball milling conditions is feasible. However, a general and broadly applicable protocol for a solvent-free mechanochemical Wittig reaction using non-stabilized phosphonium ylides and a diverse set of carbonyl compounds has not yet been established. This gap in the literature prompted us to further investigate conditions for fast and high-yielding mechanochemically driven aldehyde-to-olefin conversions. Our newly presented method is unprecedented and outstanding in terms of ease of reaction setup and reaction time, representing a significant advancement in the field of mechanochemical synthesis.

Results & Discussion

Optimization of Reaction Conditions

Throughout our optimization studies, 4-phenyl benzaldehyde (I), methyltriphenylphosphonium bromide (a, PPh₃MeBr), and a solid base were ball-milled in a 7 mL Teflon^M milling vessel containing one 12 mm stainless steel ball. The reactions were carried out at a defined frequency using an IST636 mixer mill (for detailed optimization data, see ESI).

We commenced by testing two solid bases, K_2CO_3 (1.6 equiv.) and KO^tBu (1.6 equiv.), which are frequently employed in Wittig reactions in solution, for the solvent-free methylenation of I (0.5 mmol) using PPh₃MeBr (1.4 equiv.) with a milling time of 45 minutes at 30 Hz (Figure 2, entry B7 and A1). Although previous protocols demonstrated that K_2CO_3 enables ylide formation with prolonged milling times (20 hours), this base was ineffective with a 45-minutes milling time in our protocol (entry B7).⁵⁹ Gratifyingly, employing KO^tBu as the base lead to full conversion of I, giving the desired vinyl derivative (1) in 70% isolated yield (entry A1).

Further optimization showed that the milling time could be reduced to as short as 30 seconds, with the aldehyde (I) still showing full conversion (entry A6). Consequently, 30 seconds was selected as the preferred reaction time, despite giving a lower yield compared to the other data points (entry A2- A5).





At a shorter milling time of 10 seconds, unreacted aldehyde was still present in the reaction mixture (entry A7). Without mechanical impact (0 seconds) starting material could be quantitatively recovered (entry A8). This blank experiment confirms that the reaction is not taking place during workup procedures, but that the conversion of I to $\mathbf{1}$ is indeed mechanochemically induced.

Careful finetuning of the phosphonium salt and base equivalents significantly improved the yield of the product (1) from 73% to 89% (entries B1 - B6). Additionally, variation in milling frequency showed that a threshold frequency of at least 30 Hz is necessary (entries C2 and C3); at lower frequencies, only minimal product formation was observed, with unreacted starting material being recovered (entry C1). Finally, we established a protocol that excels in ease of reaction setup and rapid reaction times (entry C3).

An olefination in solution using the same substrates and achieving a similar yield of 90% is described in the literature (Figure 1, top, left).⁶² In this conventional method, the ylide from PPh₃MeBr is pre-formed at 0 °C in THF under an inert atmosphere, using an excess of the highly pyrophoric and oxygen-sensitive base *n*-BuLi. The resulting solution is stirred for 30 minutes at 0 °C. After the addition of aldehyde I, the reaction is allowed to reach room temperature and stirred for 12 hours until complete conversion.

In contrast, our protocol involves combining all reactants - the carbonyl compound, the phosphonium salt and solid, easy-to-handle base KO^tBu - in a TeflonTM milling vessel containing one milling ball, without the need for air or moisture exclusion (Figure 1, bottom, left). The vessel is closed, mounted in a mixer mill, and ground for only 30 seconds at a frequency of 36 Hz. Typically, no pre-formation of the phosphonium ylide or sequential aldehyde addition is necessary to ensure a high-yielding olefination reaction. However, for highly base-sensitive substrates, ylide pre-formation with lower base-to-phosphonium amounts can be conducted. In this modified approach, KO^tBu (1.2 equiv.) and PPh₃MeBr (1.4 equiv.) are ground for 1 minute at 36 Hz to form a yellow paste. The vessel is then opened, the carbonyl compound is added, and milling is continued until the reaction is complete.

Notably, the progress of the reaction can be easily monitored by opening the reaction vessel multiple times to take small samples, without negatively affecting the reaction outcome. This flexibility and ease of monitoring further underscore the practicality and efficiency of our mechanochemical protocol.

Substrate Scope

With the optimized reaction conditions in hand, we evaluated the generality of this protocol by converting a diverse set of carbonyl compounds into their respective olefins (Scheme 1). Benzaldehyde derivatives (1-9, 11, and 13) consistently yielded good to excellent results, ranging from 50% to 95%, regardless of their substitution patterns. Electron-donating (2, 3, 7 - 9) and electron-withdrawing (5, 6, 7)13) groups, as well as combinations thereof (4), were well tolerated on various positions on the aromatic ring. Interestingly, under standard conditions without ylide pre-formation, product 9 was isolated in a moderate yield of 58%, with 16% of a by-product resulting from CN-sidechain dealkylation. However, using the modified procedure with ylide pre-formation (1.2 equiv. of KO^tBu and 1.4 equiv. of PPh₃MeBr), the reaction proceeded smoothly, yielding product **9** in 80% with no observable dealkylation. A pinacol boronate-substituted aldehyde, which is highly useful for subsequent Suzuki-Miyaura coupling, reacted more reluctantly. But, after a prolonged milling time of 1 hour, the desired product **11** was obtained in 80% yield. This protocol is not limited to benzaldehyde derivatives; for example, 4-methoxycinnamaldehyde reacted smoothly, even on an increased scale, yielding product 12 in 84% (0.5 mmol scale) and 80% (2 mmol scale). Garner's aldehyde, a valuable chiral building block in natural product synthesis, underwent olefination readily, yielding 80% of its vinylated analogue 16.63 Remarkably, highly conjugated all-trans-retinal was methylenated in 71% yield (product 17), underscoring the potential of this method for future industrial applications in carotenoid synthesis.²³



Scheme 1. Scope of the Wittig olefination of aldehydes and ketones using PPh₃MeBr. Reactions were performed on a 0.5 mmol scale under air in an IST636 mixer mill, using a Teflon[™] milling jar (7 mL) and one stainless steel milling ball (12 mm) at a milling frequency of 36 Hz. If not stated otherwise, milling was conducted for 30 seconds using KO'Bu (0.7 mmol, 1.4 equiv.) and PPh₃MeBr (0.6 mmol, 1.2 equiv.). Isolated yields are shown. [a] 2.4 equiv. KO'Bu were used. [b] Ylide pre-formation was conducted milling KO'Bu (1.2 equiv.) and PPh₃MeBr (1.4 equiv.) for 1 minute before aldehyde/ketone addition (for details, see SI). [c] 1 hour milling time. [d] Reaction was performed on a 2 mmol scale using two 12 mm stainless steel milling balls in a 25 mL Telfon[™] milling jar. [e] 5 minutes milling time. [f] 1 minute milling time, [g] reaction was performed on a 0.2 mmol scale.

Beyond aldehydes, ketones also performed exceptionally well under the optimized reaction conditions. Cyclohexanone-derived substrates yielded products **18** (92%) and **21** (93%). Notably, the latter is highly base-sensitive, necessitating ylide pre-formation to ensure high yields. A pyridinyl moiety was readily tolerated, and natural product (-)-menthone, along with an adamantane-derived substrate, were converted to the vinylated products **23** and **24** in 71% and 89% yield, respectively. Remarkably, the complex steroid epiandrosterone was successfully converted to product **25**, with the newly formed double bond positioned at C-17. Additionally, we demonstrated the feasibility of this approach for the late-stage olefination of the atypical antidepressant and nicotine antagonist Bupropion, yielding product **26** in 65%.^{64,65} This showcases the protocol's versatility and potential for application in the synthesis and late-stage functionalization of complex and bioactive molecules.

Subsequently, we explored the versatility of various phosphonium salts with aldehydes and ketones (Scheme 2). We selected piperonal as a model aldehyde substrate due to the benzodioxole motif's frequent presence in pharmaceutically active compounds and promising drug candidates.⁶⁶⁻⁶⁸ Generally, reactions with non-stabilized and semi-stabilized ylides (yielding products **28-33**, **37**, **38**) gave mixtures of isomers, as expected. Stabilized ylides (yielding products **34-36**) exclusively gave the *E*-isomers in yields ranging from 68% to 92%. Some substrates required slightly longer milling times due to increased steric hindrance affecting reaction rates as the substitution on the Wittig reagent increased, but no reaction was conducted for longer than 1 hour. Deuteromethylenation using PPh₃CD₃I was feasible under the given reaction conditions, yielding deuterovinylated products **27** and **39** in 81% and 90%, respectively. Remarkably, this protocol allowed for the introduction of several valuable functional groups



Scheme 2. Scope of the Wittig olefination on aldehydes I and II, and ketone III using different phosphonium halides. Reactions were performed on a 0.5 mmol scale under air in an IST636 mixer mill, using a Teflon™ milling jar (7 mL) and one stainless steel milling ball (12 mm) at a milling frequency of 36 Hz. If not stated otherwise, milling was conducted for 30 seconds using KO^rBu (0.7 mmol, 1.4 equiv.) and the respective phosphonium halide (0.6 mmol, 1.2 equiv.). Isolated yields are shown. [a] 5 minutes milling time. [b] Ylide pre-formation was conducted milling KO^rBu (1.2 equiv.) and the phosphonium halide (1.4 equiv.) for 1 minute before aldehyde/ketone addition (for details, see SI). [c] 15 minutes milling time. [d] 30 minutes milling time. [e] 1 hour milling time.

via the respective phosphonium salts: Weinreb amide (**36** and **47**), dioxolane (**37**), ester (**35** and **46**), and nitrile derivative (**38**). This versatility underscores the method's broad applicability and potential for generating a wide array of functionalized products.

Conclusion

The mechanochemical Wittig olefination protocol presented in this study offers a highly efficient and environmentally friendly alternative to conventional solution-phase methods. By eliminating the need for solvents and air- or moisture-sensitive conditions, this protocol significantly simplifies the reaction setup. The method is remarkably fast, achieving complete conversion in as little as 30 seconds, and avoids the use of highly pyrophoric bases such as *n*-BuLi. The ability to conduct the reaction without pre-forming the phosphonium ylide further underscores the practicality and efficiency of this approach.

Our results demonstrate that this protocol is broadly applicable, converting a diverse array of carbonyl compounds into their corresponding olefins with high yields. The use of different phosphonium salts to introduce various functional groups highlights the method's flexibility and potential for generating a wide array of structurally diverse products.

The simplicity and effectiveness of this mechanochemical approach make it a promising tool for advancing greener synthetic practices in both academic and industrial settings. Its rapidity, efficiency, and environmentally friendly nature offer a distinct advantage over traditional methods, contributing to a more sustainable future in synthetic chemistry.

Methods

General procedure for Wittig olefination without ylide pre-formation

A 7 mL Teflon[™] milling vessel equipped with one 12 mm stainless steel ball, was charged with the aldehyde/ketone substrate (0.5 mmol, 1 equiv.), the respective phosphonium halide (0.6 mmol, 1.2 equiv.), and potassium *tert*-butoxide (0.7 mmol, 1.4 equiv.). The closed Telfon[™] vessel was mounted into the holding station of an IST636 mixer mill and milling was conducted at a frequency of 36 Hz for 30 seconds.

After the milling process, the reaction was quenched by adding 6 mL of a sat. NH₄Cl(aq.) solution directly to the milling vessel. Subsequently, ether was added, and the contents were transferred to a separation funnel. The product was extracted three times with ether and the combined organic phases were washed once with sat. NH₄Cl_(aq.) solution, dried over Na₂SO₄, filtered, and concentrated under reduced pressure. The crude product was further purified *via* column chromatography.

General procedure for Wittig olefination with ylide pre-formation

A 7 mL Teflon[™] milling vessel equipped with one 12 mm stainless steel ball, was charged with the, the respective phosphonium halide (0.7 mmol, 1.4 equiv.), and potassium *tert*-butoxide (0.6 mmol, 1.2 equiv.). The closed Telfon[™] vessel was mounted into the holding station of an IST636 mixer mill and milling was conducted at a frequency of 36 Hz for 1 minute. Subsequently, the milling vessel was opened, and the aldehyde/ketone substrate (0.5 mmol, 1 equiv.) was added to vessel. The vessel was closed and mounted again into the mixer mill's holding station and milling process was continued at 36 Hz for additional 30 seconds.

After the milling process, the reaction was quenched by adding 6 mL of a sat. $NH_4Cl_{(aq.)}$ solution directly to the milling vessel. Subsequently, ether was added, and the contents were transferred to a separation funnel. The product was extracted three times with ether and the combined organic phases were washed once with sat. $NH_4Cl_{(aq.)}$ solution, dried over Na_2SO_4 , filtered, and concentrated under reduced pressure. The crude product was further purified *via* column chromatography.

Data Availability

The authors declare that the data supporting the findings of this study are available within the paper and its Supplementary Information.

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