Conventional methods carrying out C(sp²)–C(sp²) bond formations are typically mediated by transition-metal-based catalysts. Herein, we conceptualize a complementary avenue to access such bonds by exploiting the potential of electrochemistry in combination with organoboron chemistry. We demonstrate a transition metal-catalyst-free electrocoupaling between (hetero)aryls and alkenes through readily available alkenyl-tri(hetero)aryl borate salts (ATBs) in a stereoconvergent fashion. This unprecedented transformation was investigated theoretically and experimentally and led to a library of functionalized alkenes. The concept was then carried further and applied to the synthesis of the natural product pinosylvin and the derivatization of the steroidal dehydroepiandrosterone (DHEA) scaffold.

Despite its young history of only a few decades, the Suzuki–Miyaura reaction is one of the most utilized reactions in modern organic chemistry. The palladium-catalyzed coupling of boronic acids with organohalides was not only awarded with the Nobel prize in 2010, in fact, a recent study ranks the Suzuki–Miyaura coupling as one of the most frequently used reactions (5th place) in medicinal chemistry. Besides, many other transition-metal-mediated cross-couplings, namely Stille, Heck, Negishi, Sonogashira, Hiyama and Kumada are likewise powerful tools to forge new C–C bonds. Such indispensable strategies undoubtedly display many advantages and have inspired us to challenge the formation of C–C bonds without the need of the commonly used transition-metal catalysts, thus breaking new grounds in the field of cross-coupling reactions. We first started our ambitious concept by replacing the catalyst with an electrochemical setup. Innate advantages, including the use of inexpensive and reusable electrodes, reaction tuneability and scalability do not only rely on the modern and cutting-edge work from Baran, but trace back to many other advances in electrochemical synthesis since the pioneering works of Volta and Faraday in the 19th century.

We already employed electrochemistry to initiate aryl–aryl bond formation, inspired by the work of Geske and Waldvogel (Scheme 1A), introducing new hetero-substituted tetraarylborates (TABs). We demonstrated that the formation of unsymmetrical TAB salts is enabled by a triple ligand exchange reaction on commercially available organotrifluoroborate species employing aryl-Grignard reagents. Submitting those TABs to mild electrochemical oxidation led to the selective formation of heterocoupled biaryls (Scheme 1B).

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formation of alkenes from the corresponding alkenyl-organozinc reagents, as exemplified recently by the groups of Aggarwal and Morken (Scheme 1C). In addition, we demonstrated that the logical combination of different organometallic reagents with boron alkoxides could lead to the formation of the required bis-organoborinates in an efficient one-pot process. Based on these findings, we decided to examine the reactivity of alkenyl-aryl borate salts (ATBs) to develop an electro-olefination reaction (Scheme 1D).

ATBs (2) are underexplored salts, the only representative compound being triphenylvinyl borate which can be synthesized by treatment of tetravinyltin with triphenylborane. To investigate the electro-olefination and expand the structural variety of ATBs, we aimed to simplify their access. Therefore, we built on our previously described strategy for the synthesis of hetero-substituted tetraarylborate salts (TABS), and decided to make ATBs accessible by a triple ligand exchange reaction onto the corresponding potassium alkenyl-trifluoroborates 1 (Molander salts), employing ex situ generated Grignard or organozinc reagents.

We anticipated that the removal of an electron through an oxidation process should occur preferentially on the alkenyl moiety, avoiding the energetically disfavored deoxygenation of one of the aryl groups. As a proof of concept, we first synthesized the model systems 2a and 2b, possessing respectively para-fluorophenyl and phenyl moieties in addition to the β-styryl substituent (Figure 1). To describe the change in the electronic structure upon oxidation of 2a and 2b, spin and charge densities were computed based on Mulliken population analysis of the DFT results. Charge densities were additionally computed using the CHarges from Electrostatic Potentials using a Grid-based (ChEIPG) method. Blue areas (Figure 1A) represent positive spin densities after oxidation. Only the alkenyl substituent is selectively oxidized in both cases whereas the charge and spin densities of the other aromatic substituents only change insignificantly, confirming our assumptions.

The oxidation potentials of ATB salts 2a and 2b were determined by cyclic voltammetry and compared to the value measured for commercial sodium tetraphenyl borate (Figure 1B). With a fluoride atom present on each of the aryl groups, an Eox value of +0.81 V vs. SCE was measured for 2a, similar to the one of 2c (+0.82 V vs. SCE). However, in the absence of electron-withdrawing substituents, the oxidation potential of 2b was decreased to +0.67 V vs. SCE. As expected, it can be concluded that alkenyl groups are easier to oxidize and that the oxidation potential varies with the electronic nature of substituents on the moieties surrounding the boron atom. From a chemoselectivity perspective, the favorable oxidation of the olefin leaves no other path for the reaction but to transfer one of the remaining aryl moieties, thereby avoiding the undesirable formation of biaryl homocoupling compounds.

2a was chosen to test and optimize the reaction conditions. Inexpensive and reusable glassy carbon electrodes (GCE) proved to deliver the desired stilbene derivative 3a with optimal conversions in acetonitrile at 25°C. Following the transformation by ‘H NMR (Scheme 2) showed that the borate salt 2a is selectively oxidized into product 3a. Full conversion can be observed after 2.2 F in ‘H NMR studies and conversion-rate experiments of the electro-olefination using GC revealed that an optimal yield was obtained after 2 F. Remarkably, further oxidation resulted in consumption of the reaction product. Although no biaryl byproduct was detected in ‘H NMR, traces were found in GC. Interestingly, a third minor compound 3ab can be observed, which was identified as the epoxy-stilbene derivative of 3a. This side reaction will be discussed later with the mechanistic considerations (Scheme 7).

The synthesis of alkenyl-borate salts can be followed by 11B NMR and proved quantitative when employing either Grignard reagents or—in cases of sensitive functional groups—organozinc species.

Therefore, we started investigating the scope of the transformation using borate salts without prior purification. The reaction was first evaluated engaging (E)-alkenyltrifluoroborates 1a-g as starting materials in this two-pot sequence. Upon generation of the desired borate intermediates, those were treated with an aqueous solution to remove remaining inorganic salts and were subjected to electrochemical oxidation conditions after switching the solvent to acetonitrile. The results are depicted in Scheme 3. With electron-withdrawing substituents present on the aryl moieties, (E)-alkenes 3a-b were obtained in up to 69% yield over two steps. In the case of p-CN-substituted phenyl groups, the corresponding organozinc species had to be employed, lowering the overall yield of the 2-pot procedure (3c, 29%). This consequence decrease in yield can be attributed to the lower reactivity of organozinc deriva-
tives in ligand exchange reactions. Electron-donating and neutral aryl substituents furnished the desired (E)-alkenes in moderate to good yields (42 and 74%). Varying the substitution pattern on the alkenyl moiety did not influence the course of the reaction, and 3f–g were isolated in 55 to 71%. Heteroaryl groups were also tolerated in the electro-olefination process, furnishing structures 3h–j in up to 68% yield. Interestingly, trisubstituted double bonds also led to the corresponding olefinated aryl derivative 3k in good yield (70%). The formation of the borate aldehyde proved however difficult when an acrylate derivative was used. The introduction of 3-pyridylzinc onto a trifluoroborol acrylate and subsequent electro-olefination only gave 25% of product 3l. Notably, all derivatives were obtained with excellent (E/Z) ratios, up to 99:1.

Z-alkenyl trifluoroborates were employed next. Following the same two-pot protocol, the freshly generated Z-alkenyl-triaryl borates were engaged crude in the electro-olefination under oxidative conditions. Diverse substituted aryl moieties were able to perform the coupling reaction, furnishing compounds 3m–s in reasonable yields (43 to 64%). It is however interesting to notice that all derivatives were isolated as trans-isomers. Given that either of the starting material (E or Z) gives the same thermodynamic E isomers after electro-coupling, the strategy is stereoconvergent (Scheme 3). As it will be discussed in the mechanistic part, we assume that the oxidation of the double bond into a radical cationic species allows for the resulting bonding system to freely rotate and adopt the thermo-
dynamically more stable configuration before abstraction of the boron-containing moiety (Scheme 7).

Our study of the electro-olefination was pursued with the use of \( \alpha \)-substituted alkenyl borates (Scheme 4). The simple acyclic isopropenyl borate salt delivered product 4a in 41% yield. Cyclic alkenyl groups were then investigated in the presence of electron-rich, -neutral and -poor aromatic systems, and gave compounds 4b-f with up to 75% yield. Borate salts containing heteroatoms in the cycloalkenyl scaffolds such as 3,6-dihydro-2\( \text{H} \)-pyranyl, -thiopyranyl and 1,2,3,6-tetrahydropyridyl underwent successful electro-olefinations, delivering trisubstituted olefins 4g-o in moderate to good yields. We lastly demonstrated the reaction to be compatible in the presence of ketal functionalities (4p-r, up to 86%).

Next, we applied the method to the derivatization of more challenging structures to demonstrate the synthetic potential of our ATBs salts. Dehydroepiandrosterone (DHEA) was derivatized into a TBS-protected ether and the carbonyl function transformed into the corresponding alkenyltrifluoroborate 1o.

The addition of arylmagnesium bromide reagents to 1o, followed by electro-olefination under the optimized oxidative conditions described above furnished functionalized molecules 5a and 5b in up to 70% yield (Scheme 5A). In addition, \( \beta \)-styryltrifluoroborate 1a was employed as substrate for the synthesis of the natural product pinosylvin (Scheme 5B). 3,5-Dimethoxyphenylmagnesium bromide was introduced to perform the triple ligand exchange reaction and gave the intermediate alkenyltriazyl borate species. Subsequent electro-olefination and demethylation with BBr\(_3\) furnished 5c in 35% yield over three steps with perfect control of the diastereoselectivity \((E/Z = 99:1)\). Furthermore, the chemoselectivity was investigated on our benchmark salt 2a under distinct oxidative conditions (Scheme 5C). As already mentioned before, the electro-olefination occurs in a stereoconvergent manner. We selectively obtain the stilbene derivative 3a using \((E)\)-2a or \((Z)\)-2a in moderate to good yields. In contrast, typical Zweifel conditions led to a stereospecific inversion of the double bond configuration, as the reaction proceeds through two consecutive stereospecific steps (1,2-metallate rearrangement and anti-periplanar \( \beta \)-elimination). The \((Z)\)-isomer can therefore be synthesized using Zweifel conditions (Scheme 5C) and \((Z)\)-3a was isolated in 86% yield \((E/Z\) ratio < 1:99). Noteworthy, stereodivergent Zweifel protocols have been developed. Even though
the presented method might be less versatile than these contributions, our strategy avoids the use of highly toxic chemicals such as BrCN and PhSeCl.\(^{[16]}\)

Lastly, we set out to ascertain the mechanism of this intriguing reaction, building on conversion experiments, cyclic voltammetry and theoretical considerations (Figure 1 and Scheme 2). Crossover experiments were conducted by mixing different borate salts under electrochemical conditions, confirming the absence of products resulting from intermolecular reactions and ruling out the possibility of intramolecular processes.\(^{[14]}\)

After selective oxidation of the alkenyl moiety, a rearrangement takes place. To study the nature of this rearrangement, we synthesized borate salts containing more than a single styryl group (6a–b, Scheme 6), employing styryl-Grignard reagents as (E/Z)-mixtures, and submitted them to our electro-coupling conditions. As a reference, the desired compound 3a was obtained as the sole compound from 2a. With a salt bearing two styryl groups (6a), a product ratio of 73:27 of 3a and the diene 7 was obtained ($E/Z = 85:15$). This result points out that the transfer of a vinyl group is not preferred over the transfer of an aryl group, and therefore indicates that the rearrangement is more likely to go through a $\sigma$-bond breaking process rather than an $\pi$-addition, as for the latter an unfavorable dearomatization has to occur. Example 6b (possessing three styryl moieties) further supports this hypothesis, as 7 was obtained in 54% and 3a in 46% GC-ratio. The non-statistical distribution of products 3a and 7 in both experiments also indicates that the aryl moiety is—in such cases—a better transferable ligand than the styryl group.

In summary, the alkenyl moiety is more prone to oxidation than the aryl groups (as concluded from quantum-chemical calculations and selectivity experiments, see Figure 1 and Scheme 6) and leads to an intermediate alkyl radical cationic species [A] (Scheme 7). We then propose that further intramolecular $\pi$-addition of one of the aryl moieties undergoes a rearrangement\(^{[17]}\) towards intermediate [B] in which the C–C alkyl radical bond can freely rotate and lead to the thermodynamically favored trans product (E)-3a. Oxygen probably interacts with the reaction intermediates under formation of structure [C], as 3ab was observed in traces under air and isolated in 37% yield when the reaction was carried out under oxygen atmosphere. It is however important to note that product 3ab does not come from the oxidation of product 3a under electrochemical conditions, as confirmed by control experiments, indicating a radical pathway.\(^{[14]}\) Based on cyclic voltammetry (Figure 1), galvanostatic experiments (Scheme 2) and our findings in the previous work on biaryl electro-coupling,\(^{[7]}\) we assume that no second oxidation has to occur during the formation of the desired product 3a.

In conclusion, we have developed a new conceptual approach to alkene derivatives through electro-olefination. A simple strategy was assembled for the synthesis of alkenylborate salts (ATBs) through ligand exchanges on potassium trifluoroborates. No purification of these salts was required for the sequence to be pursued and deliver the expected coupling compounds in moderate to good yields under electrochemical oxidation. Such method represents an original and stereoconvergent alternative to the formation of functionalized olefins, opening new ways of thinking about C–C bond disconnectons.

### Scheme 6.

Electro-coupling of different mixed potassium tetraorganoborate salts. (a) In situ generated following general procedure D\(^{[6]}\) as follows for 6a: 0.5 mmol 1p and 1.0 mmol styrylmagnesium bromide. For 6b: 0.5 mmol potassium trifluoro(4-fluorophenyl)borate and 1.5 mmol styrylmagnesium bromide. (b) Product distribution ratios are determined by GC analysis on crude mixtures without isolation. Homocoupled biaryls are omitted and not included in the GC-ratios for more clarity.

### Scheme 7.

Proposed mechanism for the electro-olefination of ATB 2a.
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Conflict of interest

The authors declare no conflict of interest.

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