Addition of $H$-phosphonates to quinine-derived carbonyl compounds. An unexpected C9 phosphonate–phosphate rearrangement and tandem intramolecular piperidine elimination

Łukasz Górecki, Artur Mucha and Paweł Kafarski*

Abstract
The Abramov reaction, a base-catalyzed nucleophilic addition of dialkyl $H$-phosphonates (phosphites) to carbonyl compounds, was performed with oxidized quinine derivatives as the substrates. Homologous aldehydes obtained from the vinyl group reacted in a typical way which led to $\alpha$-hydroxyphosphonates, first reported compounds containing a direct P–C bond between the quinine carbon skeleton and a phosphorus atom. For the C9 ketones a phosphonate–phosphate rearrangement, associated with a tandem elimination of the piperidine fragment, was evidenced.

Introduction
Medicinal, organocatalytic and stereoselective properties of quinine make it the most prominent representative of *Cinchona* alkaloids [1], a group of natural compounds of a unique three-dimensional structure. The structure involves a particular arrangement of two rigid heterocyclic fragments: aromatic quinoline and chiral aliphatic quinuclidine, and a hydroxy function on the stereogenic carbon atom. Such an architecture combined with the presence of nucleophilic and electrophilic centers buried in a hydrophobic environment predestinates the molecule to asymmetric applications, such as: chiral catalysis, transition metal complexing, molecular recognition, chromatographic separation and analysis of enantiomers [2-5].

Synthetic modifications of the basic structure, motivated by an improved stereoselectivity potential of quinine, are an issue of ongoing trials [6-9]. Surprisingly, phosphorus compounds chemistry, particularity that avoiding an expansion of the core carbon skeleton [10-13], is poorly recognized and mainly
involves esterification of different phosphorus acids with the O-9-hydroxy group [14–21]. These phosphorus esters were consecutively applied in organo- and metal-assisted catalysis [14,17–20] and NMR-monitored enantiodiscrimination [21]. According to our knowledge no example of formation of a direct C–P linkage between the quinine backbone and a phosphorus atom has been reported in the literature. Stimulated by this challenge we planned to envisage a nucleophilic addition of dialkyl phosphites to quinine-based carbonyl compounds and obtain 1-hydroxyalkylphosphonate derivatives (Abramov reaction, phospho-aldol reaction [22,23]). The scope, stereochemistry and side-reactions of the addition are described.

Results and Discussion
Quinine-based carbonyl compounds were obtained by oxidation of either the secondary C9 hydroxy group to the corresponding ketone or the vinyl group to homologous aldehydes. The last-mentioned alternative demanded a protection of the OH function. This was performed via carbamoylation of quinine (1) with t-butyl isocyanate as described elsewhere (Scheme 1) [24]. A higher scale of reaction improved the yield if compared to the literature data.

Vinyl group modifications
Oxidation of the vinyl group of quinine can be carried out in two different manners to give homologous aldehydes. The one-carbon atom-shortened aldehyde 4 is the product of osmium tetroxide/periodate oxidation [25]. Depending on the reaction conditions a variable ratio of epimers at the neighboring C3 carbon atom was obtained (Scheme 2). A single-step oxidation process was not selective and produced equal amounts of diastereoisomers, 56:44 (C3 R/S, yield 95%), which was comparable to the literature data reported as 50:50 (C3 R/S, 95%) by Waddell [25] and 55:45 (C3 R/S, 80%) by Braje [26]. The two-step procedure initially involved the use of a co-oxidizer other than periodate, e.g., potassium hexacyanoferrate with catalytic amounts of osmium tetroxide to obtain the vicinal diol 3 [27]. This intermediate was preparatively separated as a 60:40 (R/S) mixture of epimers at C10. The diol compound was subsequently oxidized with NaIO4 to the aldehyde 4 with simultaneous C–C bond breakage. According to the literature an oxidative cleavage on silica in a two-phase system led predominantly to the C3 epimer of the R configuration 90:10 (yield 93%) in a short reaction time [27]. In our case, when the reaction time was prolonged to 2 hours, the overall yield remained at the same level while the diastereoselectivity was reduced to 71:29 (R/S).

The homologous aldehyde can be prepared by oxidation of the double bond in a hydroboration–oxidation sequence, however, the presence of the nitrogen atoms, particularly that of the tertiary amino group of quinuclidine, may be troublesome [28,29]. Borane complexes with heteroaromatic and aliphatic amines are considered inconveniently stable in protic solvents (water, alcohols) and dissociate only at an elevated temperature [30]. Most probably, in our case the formation of the amine–borane complex proceeded faster than the hydroboration of the vinyl group. When compound 2 was reacted with the BH3·THF complex and then oxidized with pyridinium chlorochromate (recommended PCC on SiO2 [26]) a complicated mixture of products (50% of conversion) was obtained. The mixture contained the target aldehyde 6 (minority, Scheme 2) and the corresponding alcohol (majority, ratio 1:5), both in their complexed forms (borane-tertiary amino group). Again, step-by-step approach and separation of the intermediate appeared more profitable. First, the alcohol 7 was synthesized by hydroboration of the substrate with BH3·THF under an inert atmosphere followed by oxidation of the intermediate borane 5 complex with trimethylamine oxide [31]. As the oxide also released the borane–quinuclidine complex at elevated temperature the free alcohol was obtained in a satisfactory yield. This alcohol was subjected to Swern oxidation, recommended for multifunctional compounds [32], to produce the target aldehyde 8 in 65% yield.

The obtained aldehydes 4, 6, and 8 were reacted with 1.1 equiv of diethyl phosphite. The presence of the tertiary amino group of quinuclidine was expected to be a sufficient catalytic base for the addition reaction, and furthermore to induce a diastereoselectivity. However, the expected hydroxyphosphonates were not formed, neither at room temperature after 24 hours, nor upon

![Scheme 1](image_url)
increasing the temperature to 40 °C within additional 48 hours. Addition of 0.1 equiv of Et3N initiated the reaction of 4 and 8 (Scheme 3) [33], in the case of borane complex 6 a stoichiometric amount of triethylamine (1.1 equiv) was applied.

Crude reaction mixtures were analyzed by NMR. To achieve complete separation of the 31P NMR signals and reliable assessment of the diastereomeric composition addition of 10 equiv of acetic acid was demanded. Despite the long reaction time (up to 4 days at 40 °C) the starting aldehydes were not fully consumed. Partially stereoselective addition was observed for the shorter homolog 4. The diastereomeric excess of the newly appearing stereogenic center at C10 of α-hydroxyphosphonate 9 slightly depended on the reaction conditions and the C3 absolute configuration of the substrate, and varied in the range of 40–50% (Scheme 3). The R-C3 epimer gave rise to somewhat more pronounced induction. The 31P NMR resonances of the predominating forms of the hydroxyphosphonate are shifted.
apart by approximately 1.0 ppm. We speculate that this means a
diastereomeric relationship of their C3–C10 fragment (being the
inserted spectrum in Scheme 3). Thus, general stereo-control-
lizing properties of quinine predominate and do not cooperate (no
match–mismatch effect visible) with the absolute configuration
of the starting aldehyde epimers. The hydroxyphosphonates
derived from the longer homologs were completely racemic at
C11. Two diastereoisomers of the hydroxyphosphonate
were formed in a ratio of 1:1, irrespectively of the substrate amino
group state: either free (8) or complexed with borane (6).
Elevated temperature and the presence of 1.1 equiv of Et3N
caused entire decomposition of the quinuclidine–borane com-
plex in the case of substrate 6. Final α-hydroxyphosphonate
esters 9 and 10 were purified by column chromatography and
categorized (for 9 two fractions, each containing two individ-
uals, were refined by preparative thin-layer chromatography, for
details see Supporting Information File 1).

The mixture of ketones 11 and 12 was treated with diethyl
phosphite and heated in toluene at 50 °C for a week with addi-
tion of a catalytic amount of triethylamine (Scheme 4). The
reaction mixture was purified by column chromatography. For-
mation of four diastereomeric compounds, derivatives of
eq (8R,9R), quindoline (8R,9S), quinine (8S,9R) and
equinine (8S,9S), was expected under non or partially stereo-
selective conditions. However, spectroscopic characterization
revealed the presence of only two species (one present in an
overwhelming excess) which exhibited the
NMR chemical
shifts not expected for phosphonates but typical for phosphates,
13b:
−5.76 and
−5.49 ppm. Apparently, they were products of
the phosphonate–phosphate rearrangement of intermediate
hydroxyphosphonates [36-38]. Treatment of ketones 11 and 12
with dimethyl- and diphenyl phosphite brought quite similar
results. The expected product, diphenyl hydroxyphosphonate
was not obtained, instead the quinotoxin enol diphenyl phos-
phate 13c appeared, and it was separated chromatographically
whereas methyl monodealkylated derivative 13a precipitated
directly from the reaction mixture. The selective hydrolysis of
the phosphorus esters is not surprising as triethylamine and
quinuclidine are bases strong enough to release the methyl ester.

The additional structural modifications of the quinine skeleton
of 13a were indicated with the
H,1H-C-HMBC correlation
spectra. The C2–H18 and C6–H14 interactions were present,
whereas correlations C2–H12, C6–H12, C8–H14 and C8–H18
were not visible (Scheme 5), what demonstrated a degradation
of the bicyclic fragment of quinuclidine to a piperidine skeleton.
In addition, the characteristic signal of the H11 proton was
absent and the H12 resonance was shifted to the lower field
(5.43 ppm), between the H20 and H21 vinyl protons. The C8
resonance signal was consequently shifted from 60 ppm to
approximately 120 ppm. The aromatic system remained intact.
These data suggest formation of the C8=C9 double bond in a
cascade process with concomitant cleavage of the C–N bond that follows the phosphonate–phosphate rearrangement (Scheme 6). Two $^{31}$P NMR signals are related to the E/Z diastereoisomerism. Configuration of the predominating form can be assigned as Z. First, this is indicated by the nuclear Overhauser effect – irradiation of the H12 proton caused the most significant cross-relaxation changes in intensity of the H3’ and H5’ protons of the quinoline system. This proximity is achievable only in the case of location of vinyl and quinoline protons at the same side of the double bond. Theoretical prediction of the H12 NMR chemical shift provided an additional confirmation [39]. The δ calculated for the Z arrangement (geminal alkyl, cis aromatic and trans dialkyl phosphoryl, whose estimated influence corresponds to the acetoxy group [40]) is 5.4–5.5 ppm and well matches with the observed values (5.43–5.49). The chemical shift calculation performed for the opposite configuration remains in worse agreement (5.2–5.3 ppm).
The observed reactivity seemed to be general as formation of compound 13b was evidenced (to a different extent) in other variants of the catalytical addition of diethyl phosphite to quinuclidean, with catalytic systems such as: KF/Al₂O₃, NH₃/EtOH and DBU/EtOH or toluene. Independent of the catalyst and conditions applied α-hydroxyphosphonates were not detected in the crude reaction mixture, and the rearranged compound was the only appearing product. The enol phosphates 13 were not stable and underwent slow decomposition to give four to five signals in the ³¹P NMR spectra after a month.

This is a novel contribution to the reactivity of quinine although similar eliminations of piperidine in Cinchona alkaloids have been reported in the literature. Accordingly, heating of quinine or derivatives in acids provided either quino-/cinchotoxin ketones or their tautomeric enol esters, depending on the substrate structure and the reaction conditions [41-43]. The corresponding compounds were also suggested as the products of a base-catalyzed Hofmann elimination of quaternary quinuclidinium salts studied as chiral catalysts [44,45]. These unwanted rearrangement negatively influenced the stereoselective properties of the alkaloids [44,45]. An elimination associated with the phosphonate–phosphate rearrangement was also reported for other 1-hydroxyphosphonate systems [46-48].

Conclusion
An intriguing chemical behavior of C-9 quinine-derived ketones was demonstrated in the Abramov (phospho-aldol) reaction. These carbonyl compounds reacted with dialkyl and diphenyl phosphites producing quinotoxin enol phosphates that resulted from a tandem phosphonate–phosphate rearrangement and an intramolecular piperidine elimination. It can be hypothesized that the driving force of the structural changes is the proximity of the tertiary amine nucleophilic center. Based on this supposition, a mechanism of the rearrangement was suggested. The homologous C10 and C11 aldehydes obtained by oxidation of the vinyl group reacted in a typical manner to yield α-hydroxy-phosphonates, the first described quinine-derived C–P compounds.

Supporting Information
Supporting Information File 1
Experimental and analytical data.
[http://www.beilstein-journals.org/bjoc/content/supplementary/1860-5397-10-85-S1.pdf]

Dedication
The contribution is dedicated to Prof. Roman Tyka on his 90th anniversary.

Acknowledgments
The work was financed by a statutory activity subsidy from the Polish Ministry of Science and Higher Education for the Faculty of Chemistry of Wroclaw University of Technology.

References