A de novo synthetic method to the access of N-substituted benzazepines

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Abstract: A novel, convenient procedure has been described for the construction of fluorine-containing benzazepines. The synthetic protocol starting from readily available dihydronaphthalene regioisomers is based on oxidative ring olefin bond cleavage followed by ring closure of the diformyl intermediates in the presence of some fluorine-containing primary amines across double reductive amination. The applicability of the developed synthetic method was demonstrated by the Graphical abstract:

R = benzyl, -CH2CF3, -CH2CHF2, -CH2CH2F, -CH2CH2CF3, -CH(CH3)CF3

synthesis of 13 benzazepine compounds isolated in 22–35% overall yields.

Keywords: benzazepine, fluorine, reductive amination, ring expansion, azaheterocycle, ring closing
**Introduction**

Functionalized azepanes constitute important components of several biologically relevant natural and non-natural products with interesting pharmaceutical properties. Some representatives of this class of azaheterocycles are known as antiviral agents, glycosidase inhibitors, anticancer agents or antidiabetics [1-5]. Therefore, in view of their medicinal relevance, an increasing number of synthetic methods have been described in recent years for the construction of highly substituted azepane derivatives [6-10]. Fluorine chemistry has become a rapidly expanding research area during the last 10–15 years. Because of the high impact of organofluorine molecules in drug research (approximately 25% of the drugs introduced in the market contain at least one fluorine atom) and agrochemistry, the synthesis of fluorinated organic scaffolds has been recognized to be a hot topic in synthetic organic chemistry over the past decades [11-13]. This high interest is based on the general understanding that the presence of fluorine atom(s) can influence biological property, metabolic stability, acid–base character, and lipophilicity [14-20]. Functionalized azepanes are important frameworks in small drug molecular design. Accordingly, the incorporation of F atom(s) in these seven-membered azaheterocycles has generated increasing interest in pharmaceutical research.

Benzo-fused azepines (benzazepines) including their functionalized derivatives form a relevant subclass in the area of azaheterocyclic compounds. Many representatives of these compounds, some of them found in various commercial drugs, are known to possess important biological properties. The structures of some representative drugs with a benzazepine core are presented on Figure 1.
Figure 1. Structures of some bioactive benzazepine derivatives.

For example, Lorcaserin (1) [21] is a drug used in the treatment of obesity, Ivabradine (2) [15] and Zatebradine (3) [22] are cardioprotective drugs (hyperpolarization-activated cyclic nucleotide-gated (HCN) channel blocker), Benazepril (4) [23] is known as angiotensin-converting enzyme (ACE) inhibitor, Fedovapagon (5) [23] possesses antidiuretic properties, while Fenoldopam (Corlopam) (6) [24] is a D1-like receptor agonist (Figure 1).

Despite the fact that a relatively large number of functionalized azepanes, fluorine-containing azepines, and various benzazepines with biological properties are known, there are only a very limited number of examples of fluorine-containing bioactive benzazepines available in the literature. The structure of several representatives of this group of bioactive products is collected in Figure 2 (structures 7–12) [25-27].
Considering the potential biological importance of fluorine-containing benzazepines on the one hand and the limited number of literature reports on these scaffolds on the other hand, the development of new synthetic strategies towards these structures represents a relevant challenge in synthetic organic chemistry. Within that framework, the main objective of this research involved the preparation of different types of fluorine-containing benzazepines via a convenient new approach.

**Results and Discussion**

The synthetic concept towards the construction of benzazepine scaffolds was based on our earlier findings regarding the synthesis of functionalized, saturated azaheterocyclic substances. This protocol involved the oxidative cleavage of the ring olefin bond of substituted cycloalkenes, followed by ring closing of the diformyl intermediates mediated by double reductive amination with various primary amines [28-30].

Our synthetic strategy for the creation of the benzazepine ring system started with the dihydroxylation of the ring C–C double bond of 1,2-dihyronaphthalene (1) with OsO₄ (2 mol%)/NMO in acetone at room temperature, which provided the corresponding cis-diol derivative (±)-2 [31]. Vicinal diol (±)-2 was next subjected to oxidative C–C ring cleavage with NaIO₄ in a THF/H₂O solvent system giving an unstable diformyl intermediate (I-1), which was then used further without isolation. It is well known that fluorine atoms incorporated into the structure of an organic scaffold, especially in the skeleton of an azaheterocycles, will significantly affect basic characteristics. Therefore, we intended to carry out the construction of the benzazepine skeleton by ring closing of dialdehyde intermediate I-1 using various
fluorine-containing primary amines. First we selected trifluoroethylamine as the amine component.

Scheme 1. Synthesis of benzazepine 3 containing a trifluoromethyl group.

Thus, dialdehyde I-1 resulting from the ring opening of diol (±)-2 was submitted to double reductive amination with 2,2,2-trifluoroethylamine hydrochloride in the presence of NaHCO₃ and reducing agent in CH₂Cl₂ at room temperature. After investigating various reducing agents such as NaBH₄, NaBH(OAc)₃ and NaBH₃CN, while the first two provided the desired product in low yields (12% and 18%), the reaction in the presence of the latter reagent, after 3 h, yielded the corresponding trifluoromethyl-containing benzazepine 3 in moderate yield (55%, two steps) (Scheme 1) confirming the feasibility of the proposed synthetic strategy.

In continuation, we extended the protocol described above towards the preparation of other benzazepines derivatives. Thus, diol (±)-2 was submitted to NaIO₄-mediated ring opening followed by subsequent treatment of diformyl intermediate I-1 with four different fluorine-containing primary amines: 2-fluoroethylamine, 2,2-difluoroethylamine, 3,3,3-trifluoropropylamine, and 1,1,1-trifluoropropan-2-amine. The reductive amination with the involvement of cyclization provided the corresponding fluorine-containing benzazepines 4–7 (Table 1).

<table>
<thead>
<tr>
<th>fluorine-containing amine</th>
<th>product</th>
<th>yield (two steps); compound number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compound</td>
<td>Structure</td>
<td>Yield</td>
</tr>
<tr>
<td>----------</td>
<td>-----------</td>
<td>-------</td>
</tr>
<tr>
<td>H$_2$N$__CH$_2$F</td>
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<td>H$_2$N$__CHF$_2$</td>
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<tr>
<td>H$_2$N$__CF$_3$</td>
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<tr>
<td>H$_2$N$__CF$_3$ Me</td>
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<td>43%, (7)</td>
</tr>
<tr>
<td>H$_2$N$__Ph</td>
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<tr>
<td>NH$_2$ Ph$__Me</td>
<td><img src="image" alt="Structure" /></td>
<td>47%, (9)</td>
</tr>
</tbody>
</table>

Table 1. Synthesized benzazepane derivatives 4–9 from diol (±)-2.

Obviously, this procedure could be applied for the access of non-fluorinated derivatives as well. When benzylamine or (R)-methylbenzylamine its methyl-substituted counterpart was reacted, the corresponding benzazepines 8 and 9 could be isolated in moderate yields (two steps, 47% and 55%) (Table 1).

Next, we intended to further extend the synthetic methodology and increase the number of benzazepine derivatives by targeting structural isomers. Accordingly, 1,4-dihydronaphthalene (10) a regioisomer of 1 was subjected to the same oxidative ring opening. First compound 10 under OsO$_4$-mediated dihydroxylation conditions yielded the corresponding diol (±)-11, which subsequently underwent ring opening upon treatment with NaIO$_4$ to furnish diformyl intermediate 1-2 (Scheme 2).
Scheme 2. Synthesis of benzazepine 12 containing a trifluoromethyl group.

<table>
<thead>
<tr>
<th>fluorine-containing amine</th>
<th>product</th>
<th>yield (two steps); compound number</th>
</tr>
</thead>
<tbody>
<tr>
<td>H₂N–CH₂F</td>
<td>N–CH₂F</td>
<td>25%, (13)</td>
</tr>
<tr>
<td>H₂N–CHF₂</td>
<td>N–CHF₂</td>
<td>26%, (14)</td>
</tr>
<tr>
<td>H₂N–CF₃</td>
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<tr>
<td>H₂N–CF₃ Me</td>
<td>N–CF₃ Me</td>
<td>30%, (16)</td>
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<tr>
<td>H₂N–C₆H₆</td>
<td>N–C₆H₆</td>
<td>58%, (17)</td>
</tr>
</tbody>
</table>

Table 2. Synthesized benzazepane derivatives 13–17 diol (±)-11.

Similar to Scheme 1, dialdehyde I-2 was used in the forthcoming step without isolation. Thus on treatment with 2,2,2-trifluoroethylamine in the presence of NaBH₃CN, double
reductive amination afforded by cyclization the desired benzazepine derivative 12 containing the trifluoromethyl group. This product is a regioisomer of 3 (45%, two steps) (Scheme 2).

Finally, diformyl intermediate I-2 derived from diol 11 was treated with the fluorinated primary amines 2-fluoroethylamine, 2,2-difluoroethylamine, 3,3,3-trifluoropropylamine, 1,1,1-trifluoropropan-2-amine and benzylamine under reductive condition, in the presence of NaBH₃CN, to deliver the corresponding opposite regioisomers 13–17 in moderate yield (Scheme 2).

Conclusions

In this paper we described a novel route for the construction of 2-benzazepine and 3-benzazepine ring systems starting from dihydronaphthalene regioisomers, providing a convenient access to both tetrahydrobenzo[c]azepine and tetrahydrobenzo[d]azepine regioisomers. The key steps of the synthetic procedure are (i) oxidative olefin bond cleavage of dihydronaphthalenes followed by (ii) cyclization resulting in a formal ring expansion under reductive amination with various primary amines. In view of the importance of organofluorine scaffolds, we applied fluorinated amines for the ring-closing step, which yielded various fluorine-containing benzazepines. Further extensions of the described procedure regarding the access of functionalized benzazepines are currently being studied in our laboratory.

Experimental

General procedure for dihydroxylation of dihydronaphthalene

To a solution of 1,2-dihydronaphthalene or 1,4-dihydronaphthalene (2 mmol) in acetone (30 mL) was added NMO (1.5 equiv) at 0 °C with stirring, followed by addition of a solution of 2% OsO₄ in t-butyl alcohol (0.3 mL). Next the resulting mixture was stirred for 3 h at room temperature. After termination of the reaction (monitored by TLC) 10 mL of saturated aqueous Na₂SO₃ solution was added and the reaction mixture was extracted with CH₂Cl₂ (3 × 20 mL). The combined organic layers were dried over Na₂SO₄, filtered, and evaporated under reduced pressure. The crude product was purified by means of column chromatography on silica gel (n-hexane/EtOAc or n-hexane/acetic).
was added (40 mL). The mixture was then extracted with \( \text{CH}_2\text{Cl}_2 \) (3 × 20 mL) and the combined organic layers were dried over \( \text{Na}_2\text{SO}_4 \). The resulting solution containing the dialdehyde derivative concentrated to half of its volume was used without purification for the next reaction. To the solution of the dialdehyde was added fluorine-containing amine hydrochloride (1 equiv) and \( \text{NaHCO}_3 \) (2 equiv) or benzylamine or methylbenzylamine (1 equiv, without \( \text{NaHCO}_3 \)). Then the mixture was stirred at 20 °C for 10 min and, after adding \( \text{NaBH}_3\text{CN} \) (1 equiv) and \( \text{AcOH} \) (2 drops), stirring was continued for another 3 h at 20 °C. The reaction mixture was diluted with \( \text{H}_2\text{O} \) (20 mL) and extracted with \( \text{CH}_2\text{Cl}_2 \) (3 × 20 mL). The combined organic layers were dried over \( \text{Na}_2\text{SO}_4 \) and concentrated under reduced pressure. The residue was purified by column chromatography on silica gel (\( n \)-hexane/EtOAc or \( n \)-hexane/acetone).

*Characterization and \(^1\text{H} \) NMR and \(^{13}\text{C} \) NMR spectra of the synthetized compounds are available in the Supporting Information*

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**References**


