A robust iron catalyst for the selective hydrogenation of substituted (iso)quinolones

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By applying N-doped carbon modified iron-based catalysts, the controlled hydrogenation of N-heteroarenes, especially (iso)quinolones, is achieved. Crucial for activity is the catalyst preparation by pyrolysis of a carbon-impregnated composite, obtained from iron(ii) acetate and N-arylimino-pyridinides. As demonstrated by TEM, XRD, XPS and Raman spectroscopy, the synthesized material is composed of Fe(0), Fe3C and FeN, in a N-doped carbon matrix. The decent catalytic activity of this robust and easily recyclable Fe-material allowed for the selective hydrogenation of various (iso)quinoline derivatives, even in the presence of reducible functional groups, such as nitrites, halogens, esters and amides. For a proof-of-concept, this nanostructured catalyst was implemented in the multistep synthesis of natural products and pharmaceutical lead compounds as well as modification of photoluminescent materials. As such this methodology constitutes the first heterogeneous iron-catalyzed hydrogenation of substituted (iso)quinolones with synthetic importance.

Over the last decade, implementation of low cost, less toxic and earth-abundant metals represents an attractive alternative to the use of scarce and often toxic precious metals in catalytic hydrogenation technologies. Despite substantial progress in the hydrogenation of N-heteroarenes by noble metal catalysts (e.g. Pd, Ru, Rh, Ir, Au, etc.), known homogeneous catalysts sometimes require co-catalysts/additives for substrate or catalyst activation, while heterogeneous catalysts, sometimes, lack regioselectivity and functional group tolerance. Until recently, selective hydrogenations of N-heteroarenes by base metal catalysts (e.g. Co, Ni, etc.) have been explored to a limited extent. Intriguingly, to chase sustainability and confront environmental concerns, the prime choice of any transition metal in catalysis would be utilizing iron. However, it lacks predictability and complete control over its reactivity due to competitive single electron transfer (SET), which requires sophisticated ligand systems in homogeneous catalysis.

In fact, W. D. Jones and co-workers disclosed that a molecularly defined Fe-PNP pincer complex can promote the hydrogenation of N-heteroarenes in the presence of bases (Fig. 1). In contrast to these sensitive homogeneous systems, operationally simple and robust heterogeneous iron catalysts are not known, except for one example where Shaw et al. demonstrated the hydrogenation of quinaldine at 300 °C and 5.2 MPa H₂ (Fig. 1). Recently, substantial efforts have been made to provide advanced materials with tunable structure and properties for catalysis in organic synthesis complementary to molecularly well-defined organometallic catalysts. In this context, we have developed a spectrum of supported nanostructured metal/
metal oxide particles modified by a N-doped carbon matrix and applied them in redox reactions, such as the reduction of nitroarenes, nitriles and ketones as well as the oxidation of alcohols. The synergistic combination of metal/metal oxide nanoparticles and N-doped carbon, originating from ligands upon pyrolysis, turned out to be crucial for the desired reactivity and excellent selectivity. Since nitrogen doping into the carbon matrix has a substantial influence on the catalytic activity of metal/metal oxide nanoparticles, the selection of the N-donor ligand is critical to access high reactivity and it mostly remains limited to 1,10-phenanthroline. Intriguingly, finding alternative ligands to 1,10-phenanthroline would be a significant step forward in the context of the development of catalytically active materials.

Herein, we report a series of novel supported Fe-based materials modified by a N-doped carbon matrix, generated from bidentate N-aryliminopyridine ligands upon pyrolysis. Utilization of the optimal material allowed for the first heterogeneous iron-catalyzed hydrogenation of substituted (iso)quinolones with synthetic value.

### Results and discussion

We set out our study with the preparation of a range of Fe-based materials in a sequential process: (a) modular synthesis of N-donor aryliminopyridine ligands via Schiff base condensation; (b) complexation with Fe(OAc)$_2$ (OAc = acetate); (c) deposition by wet impregnation of complexes onto suitable supports; and (d) drying and pyrolysis at variable temperatures ($500$–$900^\circ C$) based on thermogravimetric (TG) analysis (Scheme 1 and Fig. S1† respectively).

Having a library of novel Fe-based materials, we began exploring their catalytic activity in the hydrogenation of quinolines. The iron-catalyzed hydrogenation of quinolines is shown in Table 1.

| Entry | Catalyst (mol%) | Solvent | $H_2$ (MPa) | $T$ ($^\circ C$) | $t$ (h) | Conv.([yield]$^{bc}$) (%) |
|-------|----------------|---------|-------------|----------------|--------|-------------------------|
| 1     | Fe[1]/L1(3.0)@C-800 (12) | $^{1}$PrOH | 5 | 140 | 28 | 16(16) |
| 2     | Fe[1]/L2(3.0)@C-800 (12) | $^{1}$PrOH | 5 | 140 | 28 | 12(12) |
| 3     | Fe[1]/L3(3.0)@C-800 (12) | $^{1}$PrOH | 5 | 140 | 28 | 15(15) |
| 4     | Fe[1]/L4(1.5)@C-800 (12) | $^{1}$PrOH | 5 | 140 | 28 | 55(50) |
| 5     | Fe[1]/L4(1.5)@C-800 (12) | $^{1}$PrOH | 5 | 140 | 28 | 45(45) |
| 6     | Fe[1]/L4(1.5)@C-800 (12) | $^{1}$PrOH/$^{1}$H$_2$O/3/1 | 5 | 140 | 24 | 52(52) |
| 7     | Fe[1]/L4(3.0)@C-800 (12) | $^{1}$PrOH/$^{1}$H$_2$O/3/1 | 5 | 140 | 24 | 77(70) |
| 8     | Fe[1]/L4(4.5)@C-800 (12) | $^{1}$PrOH/$^{1}$H$_2$O/3/1 | 5 | 140 | 24 | 96(84) |
| 9     | Fe[1]/L4(6.0)@C-800 (12) | $^{1}$PrOH/$^{1}$H$_2$O/3/1 | 5 | 140 | 24 | 91(82) |
| 10    | Fe[1]/L4(4.5)@C-800 (12) | $^{1}$PrOH/$^{1}$H$_2$O/3/1 | 5 | 140 | 30 | >99(85) |
| 11    | Fe[1]/L4(4.5)@C-800 (12) | $^{1}$PrOH/$^{1}$H$_2$O/3/1 | 5 | 130 | 30 | 80(73) |
| 12    | Fe[1]/L4(4.5)@C-800 (12) | $^{1}$PrOH/$^{1}$H$_2$O/3/1 | 5 | 120 | 30 | 47(45) |
| 13    | Fe[1]/L4(4.5)@C-800 (12) | $^{1}$PrOH/$^{1}$H$_2$O/3/1 | 4 | 130 | 56 | >99(87) |
| 14    | Fe[1]/L4/C (~12) | $^{1}$PrOH/$^{1}$H$_2$O/3/1 | 4 | 130 | 56 | — |
| 15    | FeC@C-800 (12) | $^{1}$PrOH/$^{1}$H$_2$O/3/1 | 4 | 130 | 56 | — |
| 16    | L4@C-800 | $^{1}$PrOH/$^{1}$H$_2$O/3/1 | 4 | 130 | 56 | — |
| 17$^d$ | Fe[1]/L4(4.5)@C-800 (12) | $^{1}$PrOH/$^{1}$H$_2$O/3/1 | 4 | 130 | 56 | — |

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$^a$ Reaction conditions: $1\text{a}$ (0.2 mmol) and Fe-catalyst (12 mol% Fe) in solvent (1 mL) heated in the presence of $H_2$. $^b$ Determined by GC analysis using hexadecane as the internal standard. $^c$ GC yield reported in parentheses. $^d$ Reaction was performed in the presence of nitrogen (4 MPa).
quinolines. In an initial set of experiments with quinoline (1') in \( ^3\text{PrOH} \) in the presence of \( \text{H}_2 \) (5 MPa) at 140 °C for 28 h, the materials which were prepared from four different ligands (L1–L4) on the carbon support exhibited desired reactivity and the catalyst with L4 remained relatively more efficient (Table 1, entries 1–4). However, altering the support with L4 did not improve the conversion (Table S4, † entries 5–7).\(^8\) Interestingly, the screening of materials prepared over a wide range of pyrolysis temperatures (500–900 °C) revealed the presence of the catalyst which was prepared at 800 °C (Table S1, † entries 4 and 8–11).\(^8\) Testing polar and non-polar solvents alone did not increase product formation (Table S1, † entries 12–14),\(^8\) whereas a mixture of \( ^3\text{PrOH}/\text{H}_2\text{O} \) (3/1) enhanced the conversion slightly (Table 1, entries 5 and 6). Next, inspired by the metal-to-ligand ratio effect on the catalytic activity in the dehydrogenation of formic acid,\(^9\) we synthesized an array of catalysts with various Fe(OAc)\(_2\)/L4 ratios. To our surprise, the set of four catalysts with different Fe/L4 ratios (1 : 1.5 to 1 : 6.0) exhibited a dramatic enhancement of reaction efficiency (Table 1, entries 6–9). The progress of tetrahydroquinoline (1) formation with time resulted in the following reactivity pattern: Fe(1)/L4(4.5) @C-800 > Fe(1)/L4(6.0)@C-800 > Fe(1)/L4(3.0)@C-800 > Fe(1)/L4(1.5)@C-800 (Fig. 2). Full conversion was achieved at 150 °C and 5 MPa hydrogen pressure after 30 h (Table 1, entries 10).

Nevertheless, hydrogenation to 1 (87% GC yield) is also possible at lower temperature and pressure for a prolonged reaction time (Fe(1)/L4(4.5)@C-800 (12 mol% Fe) in \( ^3\text{PrOH}/\text{H}_2\text{O} \) (3/1), \( \text{H}_2 \) (4 MPa), 130 °C, 56 h) (Table 1, entry 13). For the success of this hydrogenation reaction, the presence of the ligand and metal as well as the pyrolysis process remains crucial. Consequently, no reactivity was observed in the absence of Fe@C-800, L4@C-800 and unpyrolyzed Fe(1)/L4(4.5)/C composites (Table 1, entries 14–16). No conversion is observed carrying out the reaction in the presence of nitrogen (4 MPa) used instead of hydrogen, which excludes transfer hydrogenation with isopropanol (Table 1, entry 17).

The composition of the catalysts was determined by elemental analysis. Interestingly, an ascending order of nitrogen amount and a descending order of iron content were noticed when the metal-to-ligand ratio was changed in the preparation process from 1 : 1.5 to 1 : 6.0 (Table S2†).\(^8\) The structural characteristics of selected materials were investigated by means of various techniques, such as powder XRD, STEM, XPS and Raman spectroscopy. Investigating the bulk phase of the materials, the powder X-ray diffraction measurements demonstrated the presence of iron carbide (Fe\(_2\)C) as a major iron constituent along with additional metallic Fe(0) and FeN\(_x\) in both the catalysts: Fe(1)/L4(4.5)@C-800 and Fe(1)/L4(1.5)@C-800, while the main components in the almost inactive Fe(1)/L4(1.5)@C-500 are iron oxides (e.g. Fe\(_3\)O\(_4\)) along with the obvious reflections for carbon in all these materials (Fig. 3 and S2†).\(^8\) Interestingly, the iron oxide (Fe\(_3\)O\(_4\)) phase was formed at lower pyrolysis temperatures (500–600 °C), while the formation of metallic Fe(0) and iron carbide (Fe\(_2\)C) was observed at higher pyrolysis temperatures (700–900 °C), likely stemming from the iron oxides (Fig. 3).\(^8\)

The electronic structure and surface elemental composition were determined by X-ray photoelectron spectroscopy (XPS) (Table S3†).\(^8\) The N1s spectra for the above-mentioned materials displayed two different peaks (Fig. S7a†). Due to similar electron binding energy (B.E.) a univocal assignment is not possible. The lower peak can be ascribed to the pyridinic nitrogen and/or nitrogen bound to the metal ion (Fe–N\(_x\)), while the higher one resembles graphitic and/or pyrrolic nitrogen.\(^21,22\) Additionally, Fe2p spectra of all samples exhibited two wide peaks centered around 711.5 and 725 eV due to 2p\(_{3/2}\) and 2p\(_{1/2}\) contributions, respectively (Fig. 4). Broadening of the peak is a hint of heterogeneity of the Fe phase present on the surface of the sample. According to TEM and literature data, this contribution is ascribed to the Fe–N bond but other Fe species in high oxidation states cannot be excluded.\(^20,22\) In addition, Fe(1)/L4(4.5)@C-800 and Fe(1)/L4(1.5)@C-800 samples clearly exhibit a contribution at a lower binding energy around 709 eV attributed to iron carbide (Fe\(_2\)C).\(^8\) The aberration corrected scanning transmission electron microscopy (STEM) high angle annular dark field (HAADF) overview images of less active Fe(1)
**Fig. 4** Fe2p XPS spectra of Fe-based materials: Fe(1)/L4(4.5)@C-800 (red), Fe(1)/L4(1.5)@C-800 (blue) and Fe(1)/L4(1.5)@C-500 (black).

L4(1.5)@C-800 (Fig. 5a) and most active Fe(1)/L4(4.5)@C-800 (Fig. 5b) catalysts were acquired together with electron energy loss spectroscopy (EELS) and energy dispersive X-ray spectroscopy (EDXS). In both catalysts, the iron containing particles, probably iron carbide according to XRD, are generally surrounded by a few to many graphene layers forming a seemingly tight enclosure. Additionally, there is a carbon phase probably stemming from the ligand, as it also contains small amounts of nitrogen (Fig. 5c and d, and S10–12). In the Fe(1)/L4(1.5)@C-800 catalyst, the additional nitrogen containing carbon structure seems to contain larger clusters of iron atoms (Fig. S12) than in the Fe(1)/L4(4.5)@C-800 catalyst, where aberration corrected high resolution STEM-HAADF images suggest the presence of finely distributed atoms and very small clusters of a heavier kind than carbon (Fig. 5e, S10 and S11†). At Fe(1)/L4(4.5)@C-800 catalyst, while they were rarely noticed in the Fe(1)/L4(1.5)@C-800 catalyst (Fig. S11 and S12†). The STEM data of almost inactive Fe(1)/L4(1.5)@C-500, prepared by pyrolysis at 500 °C, showed the parallel occurrence of nitrogen and iron in fine distribution in some regions. Also, iron oxide particles and a few iron/iron oxide core shell type particles were present (Fig. S13†). In further analysis by Raman spectroscopy, two peaks were observed at ~1590 cm⁻¹ (G band) and ~1343 cm⁻¹ (D band) in each material (Fig. S15 and Table S4†). Notably, the occurrence of the D band signified the presence of defects in the graphitic carbon matrix.

With the optimal reaction conditions in hand, we sought to explore the synthetic scope and limitations of this transformation, which are summarized in Scheme 2. A variety of quinolines with diverse substitution patterns and functionalities were employed under the developed protocol for the selective hydrogenation. At first, 2-substituted quinolines, for example, quinaldine and 2-phenylquinoline were selectively hydrogenated to corresponding 1,2,3,4-tetrahydroquinoline congeners (2), an intermediate for the synthesis of antitrypanosomal lead compounds, and 2-phenyl-1,2,3,4-tetrahydroquinoline (3), respectively, in excellent yields (both 98%). Furthermore, 3-methyl-1,2,3,4-tetrahydroquinoline (4, 90%) and 3-(4-methoxyphenyl)-1,2,3,4-tetrahydroquinoline (5, 91%) were successfully prepared from the corresponding quinoline substrates. More interestingly, a 2,3-disubstituted quinoline was hydrogenated to afford 1,2,3,4-tetrahydroquinoline derivative 6 in good yield (77%; d.r. = 1.3 : 1), whereas 2,4-dimethylquinoline was converted to 2,6-dimethyl-1,2,3,4-tetrahydroquinoline (7) in good yield (77%) and moderate diastereoselectivity (d.r. = 4.5 : 1) without arene ring hydrogenation. Aside from the pyridine ring, different substituents at the 6-position of quinoline survived under the employed hydrogenation conditions. Here, 6-isopropylquinoline was converted to 8 in moderate yield (67%). Furthermore, quinoline derivatives featuring 4-fluorophenyl and 4-trifluoromethoxyphenyl and a pyrazole moiety were successfully hydrogenated to the corresponding 1,2,3,4-tetrahydroquinoline congeners 9–11 in good to excellent yields. Gratifyingly, substituted quinoline substrates have been proven to be suitable candidates for this selective hydrogenation reaction, too. Electron-rich 6-methoxyquinoline and electron-deficient 6- and 7-fluoroquinoline were selectively converted to pyridine core hydrogenation products 12 (95%), 13 (94%) and 14 (88%), respectively, with almost equal efficiency. Furthermore, sterically hindered 8-methyl-1,2,3,4-tetrahydroquiniline (15) was also furnished in

**Fig. 5** STEM-HAADF overview images of Fe(1)/L4(1.5)@C-800 (a) and Fe(1)/L4(4.5)@C-800 (b). STEM-ADF image (c) of the lower left area of (b) overlaid with the false color image of the elemental distributions of C, N and Fe calculated from the electron energy loss spectrum imaging data set of the marked area; STEM-HAADF images with higher magnification showing details of the spectrum imaging area (d) and the extra carbon phase containing nitrogen (e) attached to the support.
very good yield (88%). In addition to the ethyl protected 8-quinolinol, which was easily converted to the reduced product 16 (92%) in excellent yield, a more challenging substrate 8-quinolinol with a free hydroxyl group was also transformed into the corresponding 1,2,3,4-tetrahydroquinolin-8-ol (28, 89%) in very good yield.

Aside from exploring the general scope, the multistep synthesis of natural products as well as medicinally important lead compounds was performed as a proof-of-applicability. Hence, the Fe-catalyzed hydrogenation was utilized as a key step in converting 2-n-pentylquinoline (30) to (-)-angustureine (31)* in two steps (Scheme 4). Furthermore, commercially available 6-methoxyquinoline (32) was transformed to 6-methoxy-1,2,3,4-tetrahydroquinoline (32, 95%), which was further converted to a tubulin polymerization inhibitor (33)* (Scheme 4).

Finally, the pharmaceutically important lead compound 37 of histamine H3-receptor antagonist was synthesized in four steps starting from readily available 6-quinolinol, which represents an attractive alternative to the noble metal catalyzed process (Scheme 4).
Additionally, the developed technology proved to be a useful tool to convert 6-pyrenequinoline-based photoluminescent materials (38 and 39) to analogous congeners (38 and 39), respectively (Fig. 6). As noticed in the UV-Vis absorption as well as emission spectra, a clear bathochromic-shift of absorption and emission wavelengths altering photophysical properties was observed due to selective hydrogenation (Fig. 6).

Since the recycling and reusability of the catalyst constitute a crucial advantage in heterogeneous catalysis, the optimal Fe-catalyst, Fe(1)/L4(4.5)@C-800, was employed seven times for the hydrogenation of quinaldine (4.0 mmol scale). To our delight, the catalyst could be recycled and reused without a significant drop of reactivity to obtain the desired 1,2,3,4-tetrahydroquinaldine (2) in almost quantitative yield (Fig. 7 and Table S5†). Notably, some Fe3O4 particles appeared in the reused catalyst compared to the fresh catalyst (Fig. S3†).

**Conclusions**

We have developed the first heterogeneous Fe-catalyst for the selective hydrogenation of quinolines and [iso]quinolones...
using molecular hydrogen as a clean reducing agent. The optimal catalyst is easily prepared from commercially available iron acetate and N-aryl iminopyridines. This robust material can be conveniently handled and is stable towards air and moisture, which provides the basis for simple catalyst recycling. A variety of quinolines with diverse substitution patterns and functionalities were successfully converted to 1,2,3,4-tetrahydroquinolines in moderate to excellent yields with high selectivity, even in the presence of other reducible functional groups, such as nitriles, halogens, esters, amides and heteroarenes. The value of this methodology is showcased by the synthesis of bioactive molecules as well as the modification of photoactive materials. It is worth mentioning that the presented Schiff base condensation of 2-pyridinecarboxaldehyde with commercially available anilines allows for the design of an infinite number of new potential catalysts for various other applications.

Conflicts of interest

There are no conflicts to declare.

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