



CrossMark
click for updates

Cite this: *Chem. Sci.*, 2015, 6, 6013

NHC-catalysed benzoin condensation – is it all down to the Breslow intermediate?†

Julia Rehbein,* Stephanie-M. Ruser and Jenny Phan

The Breslow catalytic cycle describing the benzoin condensation promoted by N-heterocyclic carbenes (NHC) as proposed in the late 1950s has since then been tried by generations of physical organic chemists. Emphasis has been laid on proving the existence of an enaminol like structure (Breslow intermediate) that explains the observed umpolung of an otherwise electrophilic aldehyde. The present study is not focusing on spectroscopic elucidation of a thiazolidene based Breslow intermediate but rather tries to clarify if this key-intermediate is indeed directly linked with the product side of the overall reaction. The here presented EPR-spectroscopic and computational data provide a fundamentally different view on how the benzoin condensation may proceed: a radical pair could be identified as a second key-intermediate that is derived from the Breslow-intermediate *via* an SET process. These results highlight the close relationship to the Cannizzaro reaction and oxidative transformations of aldehydes under NHC catalysis.

Received 17th June 2015
Accepted 20th July 2015

DOI: 10.1039/c5sc02186c

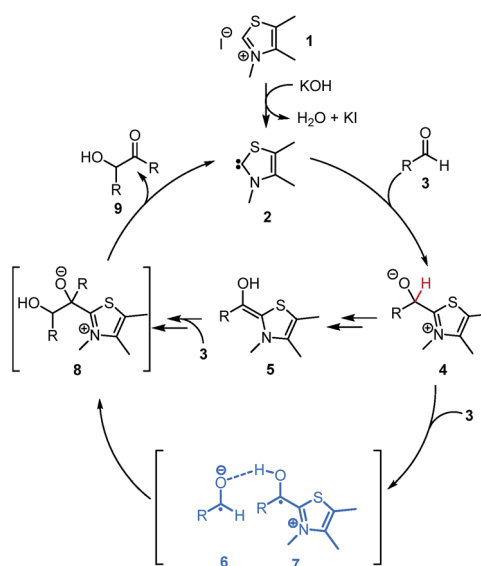
www.rsc.org/chemicalscience

Introduction

The NHC-catalysed benzoin condensation is not only of synthetic value^{1,2} but also has challenged generations of physical organic chemists with its mechanistic characteristics.^{3–5} Above all the Breslow intermediate **5** (Scheme 1) has been a long-hunted phantom which finally could be unveiled in 2012 by Berkessel's fundamental NMR studies^{5a} for a certain type of catalysts. Although one should mention that *O*-protected Breslow intermediates have been synthesized^{6b} and characterised as intermediates^{6c} of other NHC-catalysed reactions before allowing for some mechanistic interpretation. Berkessel's remarkable study on the other hand provided ¹H and ¹³C NMR data of free enaminols derived of imidazolidinium-based carbenes and aromatic aldehydes. These studies also linked the enaminol to the formation of benzoin. However, the elementary steps as depicted in the catalytic cycle (Scheme 1) still provide room for discussion: is the path of enaminol to benzoin an elementary step? How is the enaminol formed? Since for the thiazolidene catalysed benzoin condensation the only intermediate that had been characterized so far is the primary adduct **4**,^{3b} the here presented studies start from there.

Driving force for the transformation of **4** (Scheme 1) is the removal of its zwitterionic state. In principle, the charge separation in **4** can be removed by formation of the Breslow intermediate **5** (Scheme 1). The mechanism even of this simple

transformation is still not completely understood; in protic polar solvents the formal proton-shift is according to He *et al.*^{3a} most likely an intermolecular process under assistance by solvent molecules. Under aprotic conditions however, *i.e.* no Brønsted base and aprotic solvent the proton-shuffle must occur in a different fashion. Here a 1,2-hydride-shift has been discussed but could be excluded based on Berkessel's detailed studies^{5c} which could show that the resulting ketone is unreactive under benzoin condensation conditions.

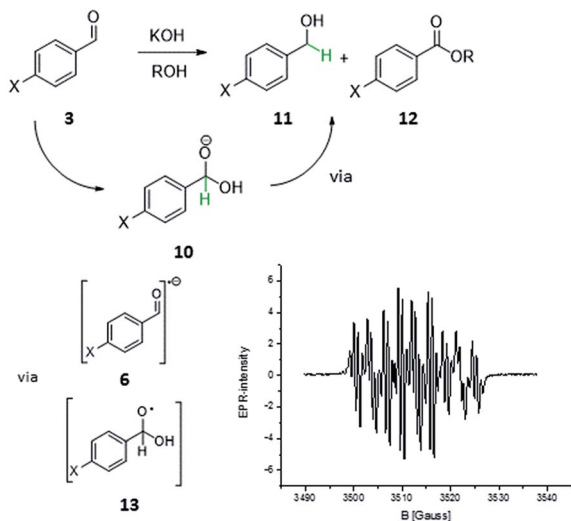


Scheme 1 Breslow catalytic cycle and a simplified alternative *via* Breslow-intermediate-like radical **7**.

Organische Chemie, Universität Hamburg, Martin-Luther-King-Platz 6, Germany.
E-mail: rehbein@chemie.uni-hamburg.de

† Electronic supplementary information (ESI) available: Characterisation data of products, substrates and catalysts, EPR and NMR spectra and progress curves as well as computational details are found. See DOI: 10.1039/c5sc02186c





Scheme 2 In analogy to the Cannizzaro process: are radical reaction steps in the formation of benzoin possible? Structurally the primary adduct **10** is closely related to **4**. The EPR spectrum was recorded of a solution of *p*-nitro benzaldehyde **3j** and KOH (1 eq.) in *i*-PrOH at room temperature.

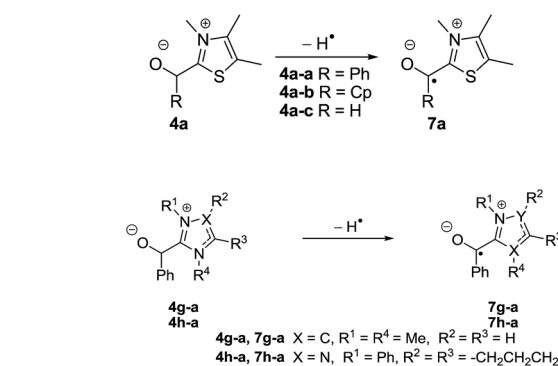
On second thoughts, one could also envision a homolytic bond cleavage of the discussed C–H bond in **4** (Scheme 1, H in red). Such a homolysis and the resulting radical are not as far-fetched as one might think. Carbon centred radicals stabilized by adjacent heterocycles^{6a,7,8,9,10,11} and heteroatoms¹² have been proposed or even characterized in several reactions that involve NHCs: the enzymatic formation of CoAsAc,⁶ the TEMPO mediated oxidative transformation of aldehydes to carboxylic acid derivatives (Studer *et al.*)⁸ and most recently the enantioselective β -hydroxylation of enals by Rovis *et al.*^{9a} and Webster, Chi and co-workers.^{9b} For understanding how the homolysis of the C–H bond of **4** within the benzoin condensation could take place it is helpful to look at the Cannizzaro reaction¹³ (Scheme 2). The primary Cannizzaro-adduct **10** of aromatic aldehyde **3** and hydroxide as base reacts with a second molecule **3** via a sequence of SET and proton transfer to benzyl alcohol **11** and benzoic acid ester **12**. This most often becomes abbreviated to a formal hydride transfer which suits our intuitive ‘polar’ understanding of reactivity. However, the ionic writing of reaction mechanisms may be just the one extreme of the whole electronic continuum of bond cleavage and formation. With a close structural relationship of **10** and **4** it is not unlikely that **4** might undergo similar SET processes.

A second advantage of the depicted radical pathway (Scheme 1) is its simplicity. The number of necessary elemental steps for the various proton shuffling events in the ionic pathway are reduced.

Results and discussion

Besides this philosophical aspect of Ockham’s razor, there is also numeric evidence. Looking at the thermodynamics by simple calculations (Table 1) the C–H bond dissociation energy (BDE) of **4** is surprisingly low, independently of the aldehyde

Table 1 Comparison of C–H BDE and RSE values ((u)B3LYP/6-31+G*) of various primary adducts **4** and radicals **7**. Values in parenthesis at (u) M06-2X/6-31+G** and in brackets (u)M06-2X/6-31+G**SCRF = PCM, solvent = 2-propanol



entry	compd	BDE [kcal mol ⁻¹]	compd	RSE ^a [kcal mol ⁻¹]
1	4a-a ^b	36.8 (39.7) [40.5]	7a-a	-50.1 (–63.9) [–54.6]
2	4a-a-H	72.0 (70.2) [69.9]	7a-a-H	-32.9 (–25.6) [–24.8]
3	4a-b	33.8 (36.7) [35.8]	7a-b	-56.0 (–66.9) [–57.8]
4	4a-c	32.6 (37.7) [38.2]	7a-c	-57.2 (–65.9) [–56.1]
5	4g-a	54.0 (47.9) [55.6]	7g-a	-59.6 (–56.9) [–59.2]
6	4h-a	38.3 (42.2)	7h-a	-66.5 (–62.6)

^a In comparison to methyl radical. ^b Designation of the compounds as follows: number of intermediate as defined in Scheme 1, first letter assigns catalyst, second letter assigns aldehyde.

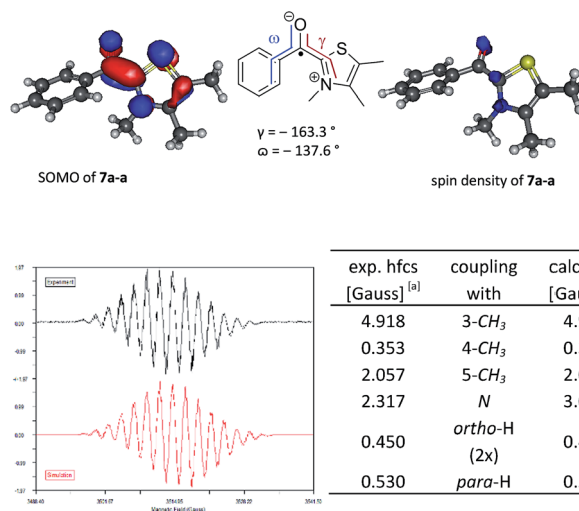


Fig. 1 Top: Plotted SOMOs and spin densities for the postulated key intermediate **7a-a** (B3LYP/6-31G*). Bottom: Experimental EPR spectrum¹⁶ and fitted EPR spectrum¹⁷ ($r^2 = 0.93$) with tabulated hyperfine coupling constants. [a] WINSIM version 0.98. [b] B3LYP/EPR-II//B3LYP/6-31G* for C, H, O, N and 6-31G* for S, Br.

should be comparably low. The high stabilization of **7** can be explained by mesomeric effects preferably by the heterocycle, since here the necessary planarity of the π -system is realized best (Fig. 1). In addition the captodative effect¹⁴ might contribute to the high stability of **7** as well.

Subsequently, typical reaction mixtures for the thiazolidene catalysed benzoin condensation in protic polar solvent (**1a**, KOH as base, *i*-PrOH, exclusion of light) and prepared under oxygen-free conditions¹⁵ were analyzed by CW-EPR spectroscopy. The observation of a strong CW-EPR-signal (Fig. 1) clearly speaks for radicals being present. Moreover, the radical is only formed if all reactants are present and air is vigorously excluded (see ESI†). The questions to be answered next are related to the structure of the radical(s) causing the EPR signal and in a second step the question of their origin and function. To address the first question we used the combination of experiments and computational data.

Led by the hypothesis that the observed EPR-signal corresponds to the suggested structures **7** or **[6·7]** one would expect that the signal should change in dependence of catalyst **2** and aldehyde **3**. Looking at the calculated spin density distribution of **7a-a** (Fig. 1 shows **7a-a**, for other examples see ESI†) one would expect the strongest dependence of the EPR spectrum on the substituents at the 3- and 5-position of the thiazole. Accordingly, the EPR-spectra resulting from the reaction solutions of 3,4,5-trimethylthiazolium iodide, KOH and various *p*-substituted benzaldehydes (**3a**, **d-f**), propionaldehyde (**3h**), and 2,2-diphenylcyclopropyl carbaldehyde (**3i**) showed only minor variations of the EPR-spectra in dependence of the aldehyde (see ESI† for EPR-spectra). On the other hand the experimental EPR-signals changed significantly as a function of the substitution pattern at the thiazolium ring (Fig. 2) supporting the postulated radical structure. Finally, calculated Fermi isotropic coupling constants (μ B3LYP/EPRIII for C, H, N, O; 6-31+G* for S) were used as starting points for fitting the observed EPR-spectra and led to fits with r^2 values ≥ 0.93 allowing the conclusion that the observed spectra actually belong to **[6·7]**. It should be noted that in several cases good fits could be obtained with simply using simulated coupling constants of **7** since the radical anions **6** often have similar coupling constants (for an EPR spectrum of **6a** see ESI†). That the EPR signal is indeed representing two radical species became apparent when analysing the EPR data derived of the reaction of **1e** and **3a**. Here the spectra could only be fitted satisfyingly if a 1 : 1 mixture of **6a** and **7e-a** were assumed (see Fig. 2 & ESI†).

Having established the structure of the experimentally observed radicals, the next step was to understand their formation, reactivity and potential role within the catalytic cycle. The simplest hypothesis for the formation of **[6·7]** is already described in Scheme 1: Another molecule aldehyde **3** acts as an oxidant of **4**. The postulated hydrogen atom transfer would benefit of a direct onwards path to benzoin **9** and free carbene **2** by simple recombination of the radical pair. Oxygen for several reasons has been excluded from being the oxidant; samples being prepared in a glove box provided EPR signals similarly to the reaction mixtures prepared under the common preparative benzoin condensation conditions (freeze-pump-

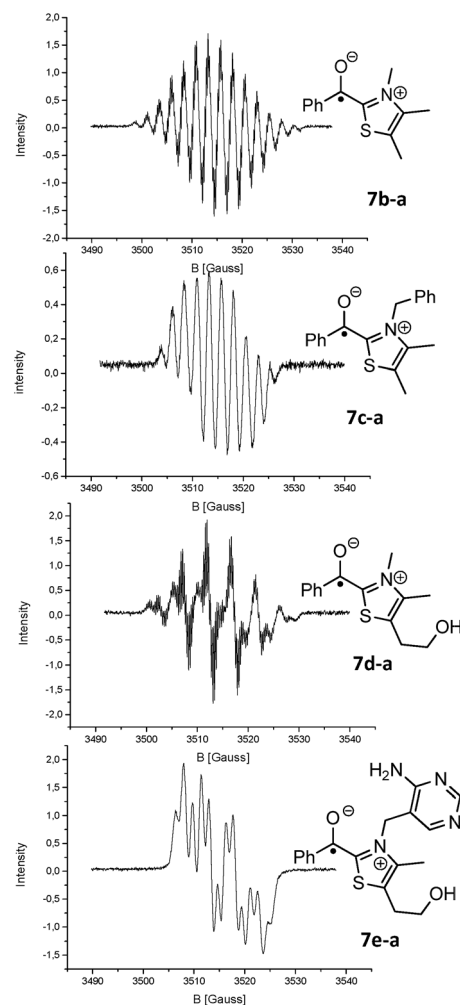


Fig. 2 EPR spectra derived of PhCHO (**3a**) in *i*-PrOH with 0.1 eq. of pre-catalysts **1b** (3,4,5-trimethyl thiazolium tetrafluoroborate), **1c** (3-benzyl-4,5-dimethyl thiazolium bromide), **1d** (3,4-dimethyl-5-(2-hydroxyethyl) thiazolium iodide) and **1e** (thiamine hydrochloride) and equimolar amounts of KOH.

thaw procedure).¹⁵ Also, opening a degassed reaction sample to air does not increase the radical signal but quenches it within 30 min and led to a different product spectrum consisting mainly of benzoic acid esters.⁸ Finally, if oxygen alone would cause the formation of **7** one would not observe a constant increase in radical with increasing catalyst load (Fig. 3) or similarly a reduction of EPR-intensity when the samples are diluted (see ESI†).

The associated progress curves (Fig. 3) are characterized by the instantaneous rise in radical concentration and a slower decline, which speaks clearly for a reactive transient. The radicals are not accumulated over time but reach an equilibrium concentration which is proportional to the initial ratio of **1** to **3**. Moreover, the shape of the progress curves also exclude a third option for the oxidant, the condensation product **9** itself. Since the highest radical concentrations are observed right in the beginning of the benzoin formation the radicals must originate from early intermediates of the catalytic cycle (Scheme 1). To follow this line of argumentation α -d₁-benzaldehyde (**3a- α -d₁**)



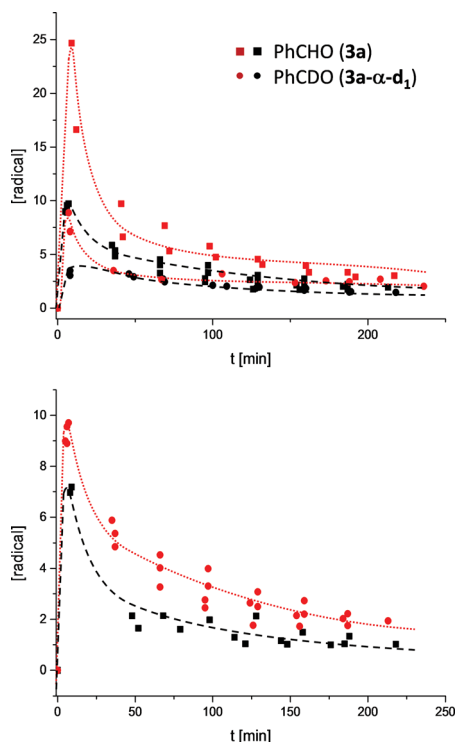
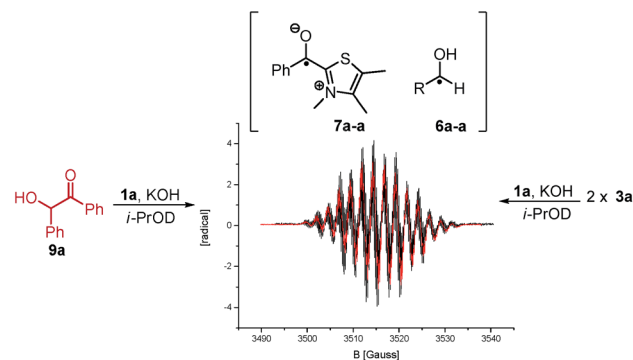


Fig. 3 Progress curves of radical under different conditions derived of 2–4 independent runs. Top: in *i*-PrOD: black symbols 0.4 eq. **1a** + KOH and 0.1 M in **3a** (square) vs. 0.1 M in **3a- α -d₁** (dot); red symbols 1.0 eq. **1a** + KOH and 0.1 M **3a** (squares) and 0.1 M in **3a- α -d₁** (dots). Below: 0.4 eq. **1a** + KOH and 0.1 M in **3a** in *i*-PrOD (red), *i*-PrOH (black).

was prepared and subjected to the standard conditions (**1a**, KOH, *i*-PrOD). If the radical is formed from the earliest, and by White and Leeper independently characterized intermediate possible,^{3b} *i.e.* **4**, the molecule needs to lose the hydrogen atom marked in red (Scheme 1). Therefore, if **7** is formed out of **4** within a sequence where the hydrogen or proton abstraction is (partially) rate determining one would expect a kinetic isotope effect (KIE).

Indeed, following the radical concentration over time for **3a** and **3a- α -d₁** (Fig. 3) a normal KIE is observed and the maximum radical concentration for **3a- α -d₁** under three different reaction conditions (see ESI†) is on average by a factor of 3.0 ± 0.7 lower than for **3a**. Keeping with Ockham's' razor, the simplest mechanistic scenario rationalizing the observed KIE would be a direct hydrogen atom transfer from **4** to **3** under the formation of **[6·7]** as depicted in Scheme 1. Interestingly, the KIE of radical formation correlates well with the observed KIE for the overall benzoin formation that has been determined by White and Leeper to be 3.4 in MeOD.^{3b} The only explanation is: benzoin and radicals must stem from mechanisms going through the same rate determining steps. This finding brings up the third question about the function of the radical. The proposed scenario in Scheme 1 describes the radical pair as a productive intermediate. The possibility that the radicals belong to a side reaction can be excluded: All tested reactions that provided an EPR signal produced benzoin and unless oxygen was present only benzoin was formed according to ¹H NMR over the



Scheme 3 Superimposed EPR spectra obtained from forward (**3a**) and reverse reaction (**9a**, 0.2 M in reactant, 0.2 eq. **1a**, 0.2 eq. KOH, *i*-PrOD). The radicals of forward and reverse reaction are identical.

course of 3.5 h. Also, based on the reversibility of the benzoin reaction and the law of microscopic reversibility further evidence was found in favour of the proposed connection radical $\leftrightarrow$ benzoin:¹⁸ Instead of **3a** benzoin **9a** was subjected to the otherwise same reaction conditions and an EPR spectrum was recorded 10 min¹⁹ after benzoin addition (Scheme 3). Not only the occurrence of a radical speaks for the link between benzoin and the proposed radical mechanism, but moreover, the obtained EPR spectrum of the reverse reaction is absolutely identical to the one derived from the forward reaction (Scheme 3).

With this information in hand we were intrigued to discover if the proposed radical pathway is an either-or-alternative to the ionic pathway or if both reaction mechanism – ionic and radical – are happening simultaneously. Therefore, conditions established by Berkessel and co-worker for forming a stable Breslow-intermediate^{5a} were used to check if **5** and **7** can be observed at the same time. Samples of **2f** and **3a** in THF-d₈ (Fig. 4) were analysed by ¹H NMR to prove that the Breslow-intermediate **5f-a** had been formed and an aliquot of the same sample was analysed by CW-EPR spectroscopy. The result is summarized in Fig. 4. Breslow intermediate **5f-a** and the corresponding radicals **[6a·7f-a]** were observed under the same conditions at the same time! In combination with the information about the kinetics and KIEs of radical and benzoin formation, this experiment provokes the question if there are indeed two simultaneous

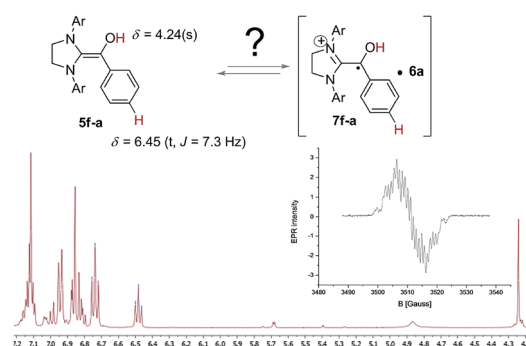


Fig. 4 Co-existence of the Breslow intermediate **5f-a** and the Breslow-type radical **7f-a** in THF-d₈.



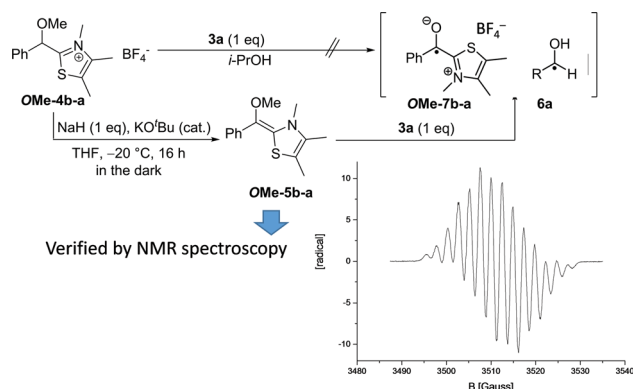


Fig. 5 Independent generation of Breslow like radicals **OMe-7b-a** from **OMe-5b-a** by addition of 1 eq. **3a**. The treatment of **OMe-4b-a** with 1eq **3a** did not provide an EPR signal indicating that at least from the cationic species **OMe-4b-a** radical formation is not directly possible *via* a hydrogen atom transfer.

pathways or if both intermediates are actually two (consecutive) intermediates of the same mechanism? The experiment in Fig. 4, however, does not allow for such a conclusion. In order to find out about a possible interconversion between 5 and [6·7] a set of experiments were carried out as described in Fig. 5.

According to a procedure of Mayr and coworkers,^{6b} *O*-methylated primary adduct **OMe-4b-a** was quantitatively transformed with stoichiometric amounts of NaH/KO^tBu (cat.) in THF into the *O*-methylated Breslow intermediate **OMe-5b-a** at −20 °C (Fig. 5). This solution was EPR silent. Only when **3a** (1 eq.) was added an EPR-signal emerged spontaneously that was by a

Table 2 Reported are peak oxidation potentials (measured against SCE in DMSO, see ref. 6a) of *O*-methylated Breslow adducts derived of **3a,d,e,f** and 3,4-dimethyl thiazole and radical concentrations and benzoin conversions determined of the same aldehydes and reaction solution of **1a** and KOH (0.1 eq.) in *i*-PrOH

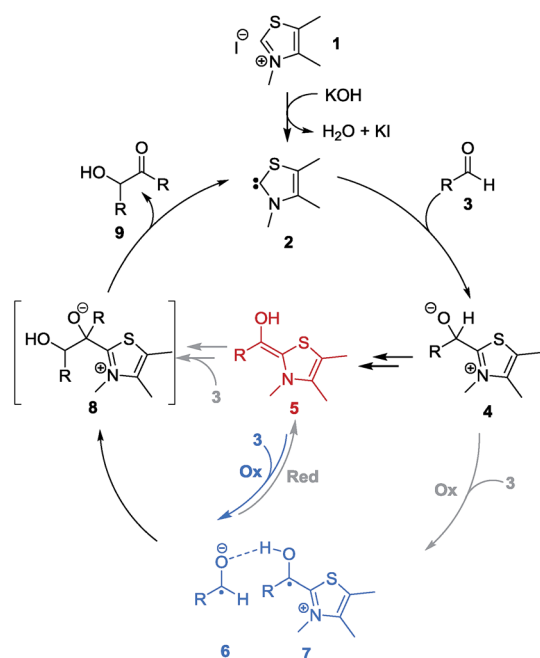
entry	RCHO	peak potential [mV]	[radical] _{max}	conv. to benzoin [%]
1	3e	−47	0.6	9
2	3d	−33	0.8	44
3	3a	−12	0.8	65
4	3f	22	1.6	80

factor of >20 more intense than any signal observed at similar substrate concentrations under normal catalysis conditions. The obtained spectrum is besides its reduced hyperfine structure²⁰ identical to the one obtained in the reaction of **3a** with **1a** and KOH to **9a**. This result finally led to a revision of the aforementioned working hypothesis (Scheme 1) and cumulates in a new catalytic cycle (Scheme 4) that incorporates the radical pair as a second key intermediate in the NHC-catalysed benzoin condensation.

Evidence that indeed the Breslow intermediate **5** is a very likely precursor of the radical pair [6·7] is found in the Table 2. Listed in column 3 are the peak oxidation potentials obtained from cyclovoltograms of **OMe-5a-a,d,e,f** in DMSO as measured by Barletta *et al.*^{6a} These values correspond very well with the maximal radical concentration observed in the NHCs catalysis of **3a,d,e,f** and **1a** (+KOH, *i*-PrOH). The oxidations potentials also correlate with the benzoin conversion obtained after 3 h. The likelihood that such a correlation is achieved by chance cannot be excluded but in combination with the so far presented results this data suggests that under the described conditions the radical pair is involved in the benzoin formation. To which extend will be clarified in future work.

Conclusions

Radical transients have been observed and characterized for a variety of N-heterocyclic carbenes and aldehydes in the benzoin condensation reaction. The structure of the observed radicals, their kinetic behavior and their instantaneous formation from benzoin under catalysis conditions provided a strong link between radicals and benzoin formation. The subsequently proven coexistence of an imidazolidinium based Breslow intermediate and its corresponding Breslow-like radical raised the question of a potential link between the proposed radical and ionic pathways. The independent generation of the same radical species as before from an *O*-methylated Breslow intermediate by addition of aldehyde answered this last question and led to the formulation of a new catalytic cycle that unites polar steps and SET–radical recombination steps. The close correlation of oxidation potentials of the *O*-methylated Breslow intermediates of four different *p*-substituted benzaldehydes with the observed radical concentrations and benzoin conversion under catalysis conditions strengthened the last conclusion. Moreover, the discovered radical intermediates can be



Scheme 4 Revised catalytic cycle – under thiazolydene-catalysis benzoin is formed most likely *via* consecutive ionic and radical steps: Blue arrow represents the newly found SET-process connecting Breslow intermediate **5** with radical pair [6·7] that collapses directly to benzoin **9** and carbene **2** *via* radical recombination.



understood as a link between other reactivities that have been exploited for instance in oxidative transformations of aldehydes *via* NHC-catalysis and also may explain the elusive nature of thiazolium based Breslow intermediates. Further studies that will provide more details on the radical formation processes and its dynamics are underway and will be published in due course.

Acknowledgements

Financial support by the FCI, DFG (Emmy-Noether-Stipend) and Fachbereich Chemie, especially C. B. W. Stark is gratefully acknowledged. We wish to thank G. Eggers for measurements of all EPR spectra presented. J. R. thanks U.-P. Apfel (Universität Bochum) for a generous donation of TritSNO.

Notes and references

- Review articles: (a) R. Kluger and K. Tittmann, *Chem. Rev.*, 2008, **108**, 1797; (b) N. Marion, S. Diez-Gonzalez and S. P. Nolan, *Angew. Chem.*, 2007, **119**, 3046; *Angew. Chem., Int. Ed.*, 2007, **26**, 2988. Latest Reviews concerning method development and synthetic applications: (c) S. de Sarkar, A. Biswas, R. C. Samanta and A. Studer, *Chem.-Eur. J.*, 2013, **19**, 4664; (d) A. Grossmann and D. Enders, *Angew. Chem.*, 2012, **124**, 320; *Angew. Chem., Int. Ed.*, 2012, **51**, 314; (e) X. Bugaut and F. Glorius, *Chem. Soc. Rev.*, 2012, **41**, 3511; (f) T. Dröge and F. Glorius, *Angew. Chem.*, 2010, **122**, 7094; *Angew. Chem., Int. Ed.*, 2010, **49**, 6940; (g) J. Douglas, G. Churchill and A. D. Smith, *Synthesis*, 2012, 2295; (h) K. Zeitler, *Angew. Chem.*, 2005, **117**, 7674; *Angew. Chem., Int. Ed.*, 2005, **44**, 7506.
- Organocatalysis, selected examples: (a) A. Grossmann and D. Enders, *Angew. Chem.*, 2012, **124**, 320; *Angew. Chem., Int. Ed.*, 2012, **51**, 314; (b) D. Enders and A. A. Narine, *J. Org. Chem.*, 2008, **73**, 7857; (c) D. Enders, O. Niemeier and A. Henseler, *Chem. Rev.*, 2007, **107**, 5606; (d) F. Glorius and C. Burstein, *Angew. Chem.*, 2004, **116**, 6331; *Angew. Chem., Int. Ed.*, 2004, **43**, 6205.
- (a) Y. He and Y. Xue, *J. Phys. Chem. A*, 2011, **115**, 1408; (b) M. J. White and F. J. Leeper, *J. Org. Chem.*, 2001, **66**, 5124; (c) G. L. Barletta, Y. Zou, W. P. Huskey and F. Jordan, *J. Am. Chem. Soc.*, 1997, **119**, 2356; (d) Y.-T. Chen, G. L. Barletta, K. Haghighi, J. T. Cheng and F. Jordan, *J. Org. Chem.*, 1994, **59**, 7714.
- (a) R. Breslow, *J. Am. Chem. Soc.*, 1957, **79**, 1762; (b) R. Breslow, *J. Am. Chem. Soc.*, 1958, **80**, 3719.
- (a) A. Berkessel, V. R. Yatham, S. Elfert and J.-M. Neudörfl, *Angew. Chem.*, 2013, **49**, 12537; *Angew. Chem., Int. Ed.*, 2013, **52**, 11158; (b) A. Berkessel, S. Elfert, V. R. Yatham, J.-M. Neudörfl, N. E. Schlörer and J. H. Teles, *Angew. Chem.*, 2012, **124**, 12537; *Angew. Chem., Int. Ed.*, 2012, **51**, 12370; (c) A. Berkessel, S. Elfert, K. Etzenbach-Effers and J. H. Teles, *Angew. Chem.*, 2010, **49**, 7120.
- (a) G. Barletta, A. C. Chung, C. B. Rios, F. Jordan and M. Schlegel, *J. Am. Chem. Soc.*, 1990, **112**, 8144; (b) B. Maji and H. Mayr, *Angew. Chem., Int. Ed.*, 2012, **51**, 10408; (c) V. Nair, S. Bindu, V. Sreekumar and N. P. Rath, *Org. Lett.*, 2003, **5**, 665.
- (a) E. Chabriere, X. Vernede, B. Guigliarelli, M. H. Charon, E. C. Hatchikian and J. C. Fontecilla-Camps, *Science*, 2001, **294**, 2559; (b) S. O. Mansoorabadi, J. Seravalli, C. Furdul, V. Krymov, G. J. Gerfen, T. P. Begley, J. Melnick, S. W. Ragsdale and G. H. Reed, *Biochemistry*, 2006, **45**, 7122.
- J. Guin, S. de Sarkar, S. Grimme and A. Studer, *Angew. Chem.*, 2008, **120**, 8855; *Angew. Chem., Int. Ed.*, 2008, **47**, 8727.
- (a) N. A. White and T. Rovis, *J. Am. Chem. Soc.*, 2014, **136**, 14674; (b) Y. Zhang, Y. Du, Z. Huang, J. Xu, X. Wu, Y. Wang, M. Wang, S. Yang, R. D. Webster and Y. Robin, *J. Am. Chem. Soc.*, 2015, **137**, 2416.
- Y. Du, Y. Wang, X. Li, Y. Shao, G. Li, R. D. Webster and Y. R. Chi, *Org. Lett.*, 2014, **16**, 5678.
- A. I. Vovk, A. F. Babicheva, V. D. Pokhodenko and A. A. Yasnikov, *Dopov. Akad. Nauk Ukr. RSR, Ser. B: Geol., Khim. Biol. Nauki*, 1977, **10**, 907.
- Heteroatoms in general are able to stabilize (carbon-centered) radicals by interaction of their free electron pairs with the half-filled p-orbital of the radical center.
- (a) E. C. Ashby, D. T. Coleman III and M. Gamasa, *Tetrahedron Lett.*, 1983, **24**, 851; (b) A. Fuentes and J. V. Sinisterra, *Tetrahedron Lett.*, 1986, **27**, 2967; (c) A. Fuentes, J. M. Marinas and J. V. Sinisterra, *Tetrahedron Lett.*, 1987, **28**, 2947; (d) E. C. Ashby, D. Coleman and M. Gamasa, *J. Org. Chem.*, 1987, **52**, 4075.
- H. Viehe, Z. Janousek, R. Merényi and L. Stella, *Acc. Chem. Res.*, 1985, **18**, 148.
- The standard conditions for sample preparation used up to five cycles of the freeze-pump-thaw-procedure (under fine vacuum). Preparation in a glove box was done for matters of comparison for a couple of samples (O_2 -content < 20 ppm).
- Reaction conditions: 0.4 M **3a**, 0.1 eq. **1a**, 0.1 eq. KOH in *i*-PrOH; presented spectrum recorded 40 min after addition of **3a**. EPR spectrometer parameters: $\nu = 9.859$ GHz, field resolution 0.049, 10.5 mW, $H_0 = 3513$ G.
- Based on calculated spin densities and Fermi contact couplings (B3LYP/EPR-III//B3LYP/6-31G*) for C, H, O, N and 6-31G* for S, Br. Fitting was done with software WINSIM version 0.98; D. R. Duling, *J. Magn. Reson., Ser. B*, 1994, **104**, 105.
- Intermediate **8** is not stable, as it is residing on a flat plateau with no significant barriers to the forward or reverse reaction as calculations by He *et al.* could show (see citation **3a**). Therefore, one can formulate a direct elemental step involving **9** and [6·7].
- Time delay due to necessary adjustments of the spectrometer.
- The lack of hyperfine structure in the spectrum is caused by the high radical concentration and potentially also by the solvent. The radical concentration was estimated to be in the region of 1.5 mM. Estimation was done using a calibration curve with TEMPO in toluene (ESI,† A.2.2.9).

