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Ruthenium-Catalyzed Hydroformylation of the Functional Unsaturated Fatty Nitrile 10-Undecenitrile

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Graphical abstract

Abstract

The hydroformylation of 10-undecenitrile (**1**), a route towards polyamide-12, has been studied using Ru-diphosphite catalysts. The reactions proceeded effectively by *in situ* combination of chloro precursors such as RuCl₂(PPh₃)₃ and RuCl₂(DMSO)₄ with Biphephos. High productivities (TON up to 15,000 mol(aldehyde).mol(Ru)⁻¹) were achieved by carrying the reactions at low catalyst loading ([**1**]₀/[Ru] = 20,000), at 120 °C in toluene or acetonitrile under 20 bars CO/H₂ (1:1), with 20 equiv of Biphephos vs. Ru. Up to 75% chemoselectivity for the aldehydes and very high regioselectivities for the linear aldehyde (*l/b* = 99:1) were reached under such optimized conditions. Lower loadings of Biphephos (down to 2.5 equiv. vs. Ru) did not affect the chemo- and regioselectivities but the activity. The Ru-Biphephos combinations showed a non-optimized hydroformylation TOF_{HF} of *ca.* 2-7 min⁻¹, that is *ca.* 1-2 order of magnitude lower than that of analogous Rh-based systems (TOF_{HF} = *ca.* 80 min⁻¹). These Ru-Biphephos systems are, however, incapable, under the conditions suitable

for selective hydroformylation, to promote isomerization of internal olefins, and hence to achieve a tandem isomerization-hydroformylation process.

Keywords: Hydroformylation, Isomerization, Ruthenium, Biphephos, 10-Undecenitrile.

1. Introduction

If rhodium is irrefutably the most efficient metal to promote olefin hydroformylation, one of the most widely applied homogeneously-catalyzed processes in industry [1], its very high and volatile price has urged investigation on other metals [2]. In 1977, the relative activities of the unmodified metal carbonyl complexes in hydroformylation were suggested as follows: Rh $\gg$ Co > Ir, Ru > Os > Pt > Pd $\gg$ Fe > Ni [3]. However, recent reports have shown that those old assumptions should be re-examined; for instance, the activity ratio of rhodium-to-iridium is in fact much closer to 1 than the 10,000:1 ratio initially predicted [4,5]. Ruthenium may also offer an interesting compromise between price and activity, as it is currently *ca.* 15 and 12 times cheaper than rhodium and iridium, respectively [6], and its activity in an oxo process is generally announced as one of the best (with iridium) among all alternative metals.

The first investigations on Ru-catalyzed hydroformylation began as early as in 1965 with Wilkinson's brief report on hydroformylation of 1-pentene using the mononuclear zerovalent complex $\text{Ru}(\text{CO})_3(\text{PPh}_3)_2$ as catalyst precursor (100–120 °C, 100 bar, $\text{CO}/\text{H}_2 = 1:1$) [7]. The authors subsequently presented more detailed results for 1-hexene hydroformylation with the same catalyst system and other related mononuclear Ru-phosphine complexes; at a quite high catalyst loading ($[\text{olefin}]/[\text{Ru}] = 100$), the turnover frequency (TOF) reached 0.075 min^{-1} , with a claimed 100% chemoselectivity for the aldehydes in most cases, although the linear-to-branched ratio was low ($l/b = 2.0\text{--}2.9$) [8]. The Ru(II) dihydrido

dicarbonyl complex $\text{Ru}(\text{H})_2(\text{CO})_2(\text{PPh}_3)_2$ was proposed as the principal active species. $\text{Ru}_3(\text{CO})_{12}$ proved to be a modest precursor (24% conv.) under the studied conditions but increased conversion was obtained upon addition of 1 equiv. (vs. Ru) of PPh_3 (88% conv.) or, even better, $\text{P}(\text{OPh})_3$ (95% conv.); however, those systems were all less active than the mononuclear complexes. Meanwhile, Schulz and Bellstedt also reported hydroformylation of propylene with $\text{Ru}_3(\text{CO})_{12}$ to afford 94% of conversion, but the final mixture contained less than 25% of aldehydes [9].

Examples of 1-hexene hydroformylation conducted in an ethanol-water (80:20) mixture using water-soluble complexes of the type $\text{K}[\text{Ru}(\text{EDTA-H})\text{Cl}]$ were reported in 1988 [10]. At a high catalyst loading ($[\text{olefin}]/[\text{Ru}] = 140$; 130 °C, 50 bar CO/H_2 1:1), these led to full conversion of the olefin (overall TOF = 0.2 min^{-1}) and exclusive formation of linear heptanal.

Surprisingly, ruthenium-catalyzed hydroformylation with the ligands most often used nowadays in combination with rhodium, *i.e.* diphosphines and diphosphites, was not reported until recently. It is only in 2012 that Nozaki and coworkers reported on combinations of $\{\text{RuCp}(\text{acac})\}_2$ with Xantphos or Bisbi diphosphines, or the A4N3 diphosphite (Figure 1) [11]. The latter diphosphite ligand allowed reaching increased chemo- (up to 66% aldehydes) and regioselectivities (*l/b* up to 79) in the hydroformylation of 1-decene (100 °C, 20 bar CO/H_2 1:1); the side-products were essentially isomerized (internal) olefins (19%) and a slight amount of the hydrogenation product (1.5%). The catalyst loading was, however, quite high ($[\text{olefin}]/[\text{diphosphite}]/[\text{Ru}] = 40:2:1$) and overall TOFs were about 0.025 min^{-1} .

Domino hydroformylation-hydrogenation reactions, to end up with the corresponding alcohols instead of the aldehydes, were also developed. Besides examples relying on rhodium complexes to achieve hydroformylation and ruthenium complexes for the hydrogenation

reaction [11,12], Beller and coworkers developed the first such domino reaction with the same ruthenium catalyst. Using $\text{Ru}_3(\text{CO})_{12}$ or $\text{Ru}(\text{methylallyl})_2(\text{COD})$ as precursor, combined with 1 equiv. (vs. Ru) of a 2-phosphino-substituted imidazole ligand, at a [olefin]/[Ru] ratio of 167, 130 °C and 60 bar CO/H_2 (1:5), they achieved full conversion of 1-octene (overall TOF = 0.13 min^{-1}) to provide 87% of alcohol ($l/b = 10$), along with 9% of octane and less than 1% of the intermediate aldehyde [13,14].

In previous studies, we reported the use of Rh-Biphephos [15] and Ir-Biphephos [5c] catalyst systems for the tandem isomerization-hydroformylation [16,17] of the unsaturated fatty nitrile 10-undecenitrile (**1**) (Scheme 1), as a route toward biosourced polyamide-12. Those systems performed at very high substrate-to-catalyst ratio (20,000–100,000) and yielded the desired linear aldehyde (**2**) with high chemo- and regioselectivities up to 93% and 99%, respectively. However, significant amounts of undesired isomerization products (**1-int-x**) along with minute amounts of the hydrogenation product (**4**) were formed, which eventually plague both conversions and selectivities for the desired linear aldehydes (Scheme 1). Preliminary experiments showed that potentially interesting results could be also reached with ruthenium catalysts, although the activities were apparently much lower [5c]. Herein we report full details on the isomerization-hydroformylation of 10-undecenitrile with such ruthenium-based systems. A variety of precursors and ligands, as well as regular reaction parameters (solvent, temperature, syngas pressure, substrate concentration), have been screened.

2. Experimental Section

2.1. General features

All reactions involving Ru-phosphine catalysts were performed under an inert atmosphere (argon) using standard Schlenk techniques. Solvents (toluene, THF, etc.) were purified over alumina columns using a MBraun system. $\text{RuCl}_2(\text{PPh}_3)_3$, $\text{RuCl}_2(p\text{-cymene})$ and $\text{Ru}_3(\text{CO})_{12}$ were generously provided by Umicore Co and stored in the glove box. $\text{RuCl}_2(\text{DMSO})_4$ was synthesized according to the literature procedure.[18] Biphephos and A4N3 diphosphite ligands were purchased from Strem Chemicals and MCAT, respectively, and used as received (stored in the glove box). 10-Undecenitrile (typically 94% pure, contains 6% of 9-undecenitrile (**1-int-0**) and other internal isomers (**1-int-x**), as determined by NMR) was supplied by Arkema; it was first eluted through a short alumina column and vacuum-distilled (Kügelrohr distillation) at 125 °C under 0.03 mm Hg prior to use. ^1H and ^{13}C NMR spectra were recorded on Bruker AC-300 and AM-400 spectrometers. ^1H and ^{13}C chemical shifts were determined using residual signals of the deuterated solvents and were calibrated vs. SiMe_4 .

2.2. General Procedure for Hydroformylation Reaction

In a typical experiment, the ruthenium precursor $\text{RuCl}_2(\text{PPh}_3)_3$, as a 1.0 g.L^{-1} toluene solution (0.72 mL, 0.75 μmol) was added on Biphephos (11.8 mg, 15.0 μmol) in a Schlenk flask. 10-undecenitrile (2.479 g, 15.0 mmol) in the desired solvent (toluene or acetonitrile, 15 mL) was added onto the resulting mixture. The solution was transferred under argon into a 90 mL stainless-steel autoclave under argon, equipped with a magnetic stir bar cross. The reactor was sealed, charged with CO/H_2 at the desired pressure at room temperature, stirred (800 rpm) and then heated with silicon oil set at the desired temperature. During the reaction, aliquots were sampled at regular time intervals to monitor the conversion and selectivities by NMR. After the appropriate reaction time, the reactor was cooled to room temperature and vented to atmospheric pressure. The solution was analyzed by NMR (after evaporation of

solvent). The conversion of **1** into **1-int-x** and **2–5**, as reported in Tables 1-5, was calculated taking into account the quantity of internal isomers (**1-int-x**) initially present in the substrate: $\text{Conv}(\mathbf{1}) = ([\mathbf{2}]_t + [\mathbf{3}]_t + [\mathbf{4}]_t + [\mathbf{5}]_t + [\mathbf{1-int-x}]_t - [\mathbf{1-int-x}]_0) / [\mathbf{1}]_0$. The reported TOF values are overall values calculated from the conversion at total reaction time: $\text{TOF} = \text{conv} \times 20,000 / \text{time}$.

The NMR characteristics for 10-undecenitrile (**1**), its internal isomers (**1-int-x**), the hydroformylation products (**2** and **3**) and the hydrogenation product (**4**) have been reported previously [5c,15]. Typical ^1H NMR signals for the linear alcohol (**5**) were observed at $\delta = 3.62$ (t, $J = 6$ Hz, 3H, $\text{HOCH}_2\text{CH}_2-$) ppm.

3. Results and Discussion

First hydroformylation experiments were performed at low catalyst loading ($[\mathbf{1}]_0/[\text{Ru}] = 20,000$; initial **1/1-int-x** ratio = 94:6) using different chloro Ru(II) ($\text{RuCl}_2(\text{PPh}_3)$, $\text{RuCl}_2(p\text{-cymene})$, $\text{RuCl}_2(\text{DMSO})_4$) and Cl-free Ru(0) ($\text{Ru}_3(\text{CO})_{12}$) precursors in combination with Biphephos, diphosphite A4N3 or triphenylphosphite. For the sake of comparison, the experimental conditions used were those optimized in the hydroformylation of **1** using $\text{Rh}(\text{acac})(\text{CO})_2\text{-Biphephos}$ [15].

3.1. Preliminary Notes

It is here important to point out that the chemo-/regioselectivities remained constant over time, and no obvious change in the kinetic regime was noted, indicating the stability of all these catalytic systems over the reaction course. This indicates, in particular, that these Ru catalyst systems were not affected by the HCl released from the chloro precursors, which is in contrast with the highly sensitive Rh-based systems. Also, all the experiments reported in the forthcoming tables were at least duplicated, showing a good reproducibility. One necessary

condition is required to achieve such reproducibility, that is the use of freshly prepared (*i.e.*, no older than one week) stock solutions of the Ru precursor, stored in the dark. Because of the low catalyst loading, such stock solutions were needed to introduce accurately small amounts of the Ru precursor. These stock solutions are perfectly clear ($\text{RuCl}_2(\text{PPh}_3)_3$: orange, $\text{RuCl}_2(\text{DMSO})_4$: yellow); yet, we observed that over days-weeks, upon exposure to light, some of these solutions can turn blackish, suggesting the formation of metal colloids/nanoparticles; in those cases, we observed that the chemoselectivity for aldehydes can dramatically drop from 75% down to 60% (mainly in favor of hydrogenation and also isomerization products), while the regioselectivity slightly dropped from 99.0:1.0 down to 98.5:1.5. We also noted that old, still clear solutions to the naked eye can lead to decreased selectivities.

3.2. Catalyst Precursor

Significant differences in terms of activity were noted among the four precursors investigated. Representative results are summarized in Table 1. Both $\text{RuCl}_2(\text{PPh}_3)_3$ and $\text{RuCl}_2(\text{DMSO})_4$ led to the higher activities with, respectively, important and full conversion of the substrate under the chosen conditions ($\text{TOF}_{\text{HF}} = 5$ and 6 min^{-1} , entries 1 and 4, respectively). On the other hand, the chemoselectivity for aldehydes was very similar for all systems. This selectivity is quite comparable to that obtained with the Rh-Biphephos catalytic system [15], so as the regioselectivity (**2/3**; *l/b* up to *ca.* 120); only the system based on $\text{RuCl}_2(p\text{-cymene})$ was somewhat less regioselective. The close similarity of the regioselectivity achieved with Cl-containing and Cl-free ($\text{Ru}_3(\text{CO})_{12}$) precursors is noteworthy (*vide supra*). The **1-int-0/1-int-x** ($x > 0$) ratio indicates the isomerization ability of the system: the lower this ratio, the more important the isomerization of the double bond inside the carbon chain. In fact, those Ru systems proved moderately isomerizing, since most of them led essentially to 9-undecenitrile

with quite minor amounts of more internal isomers. A consequence of this low isomerizing ability is that, even upon long exposure (75 h) after full conversion of **1**, the amount of internal isomers **1-int-x** did not decrease significantly. The final amount of hydrogenated product (**4**) is similar in Ru systems as in the Rh ones (*ca.* 2–5%). Noteworthy, in contrast to Rh-based catalysts but in line with Ru systems [10–14], small amounts (<0.5 mol%) of alcohol products (**5**)¹⁹ can be observed. Further experiments were conducted using the two most efficient precursors, *i.e.*, RuCl₂(PPh₃)₃ and RuCl₂(DMSO)₄.

3.3. Solvent

The dependence of the performance of the RuCl₂(PPh₃)₃- and RuCl₂(DMSO)₄-Biphephos systems on the solvent nature was evaluated with toluene, acetonitrile, DMF, diglyme and 1,2-dichloroethane (DCE). The results are summarized in Table 2. The trends observed with RuCl₂(PPh₃)₃ are identical to those with RuCl₂(DMSO)₄. Hence, acetonitrile, DMF and toluene featured similar results in terms of conversion of **1** and chemoselectivity for aldehydes. Nevertheless, if the regioselectivity was as good as the one obtained with toluene for acetonitrile, DMF induced a slightly more important amount of branched aldehyde. On the other hand, both 1,2-dichloroethane and diglyme led to low substrate conversion and also a slightly lower regioselectivity. There is no apparent correlation between the solvent polarity and the catalytic performance. Hence, this screening allowed us to highlight two couples of efficient precursors –RuCl₂(PPh₃)₃ and RuCl₂(DMSO)₄– and solvents–toluene and acetonitrile.

3.4. Ligands

The A4N3 ligand (Figure 1) used by Nozaki *et al.* [11] and triphenylphosphite, a simple monophosphite ligand, were evaluated in comparison with Biphephos, in combination with RuCl₂(PPh₃)₃ (similar results were obtained upon using RuCl₂(DMSO)₄) under the same reactions conditions than the standard experiments presented above; the amounts of ligands

were adjusted to match the same phosphorus-to-metal ratio. Unsurprisingly, the P(OPh)_3 -based system exhibited a lower activity and chemoselectivity and a much poorer regioselectivity ($l/b = 4.5$) as compared to the two other systems based on diphosphite ligands. The A4N3 ligand exhibited a slightly lower activity ($\text{TOF} = 3\text{--}5 \text{ min}^{-1}$) and also a slightly decreased regioselectivity than the equivalent Ru-Biphephos system. The same trend was also observed in the case of Rh-based catalysts in previous studies carried in our lab [20]. On the other hand, very poor performances were observed with Zhang's tetraphosphine [21].²²

3.5. *[Ligand]/[Ru] ratio*

In our standard conditions, a ligand-to-Ru ratio of 20 is used. In the case of Rh-Biphephos catalysis, this condition proved to be necessary to prevent catalyst decay and decreased activities and selectivities, assumed to arise from the formation of rhodium aggregates [15]. With ruthenium, the amount of ligand introduced may not need to be necessary as high as in the case of rhodium to prevent such phenomenon. Thus, experiments at $[\text{Biphephos}]/[\text{Ru}]$ ratios in the range 0–20 were carried out; the results are summarized in Table 4. Experiments conducted without Biphephos ligand proved to be inefficient to perform hydroformylation of undecenitrile in 88 h; at the end of these experiments, a black suspension was recovered, suggesting the formation of ruthenium aggregates. Formation of these aggregates was prevented by stabilization of the metal center with the diphosphite ligand. In these experiments, no impact was observed regarding the selectivities, *i.e.*, the l/b ratio and %HF did not change significantly when the $[\text{Biphephos}]/[\text{Ru}]$ was modified, indicating that the same active species is at work in all cases. However, a noticeable increase in the catalytic activity was noted when the $[\text{Biphephos}]/[\text{Ru}]$ ratio increased. Although this may appear counterintuitive at first sight, this observation can be rationalized by taking into account that, at such a high substrate-to-catalyst ratio (20,000), excess ligand induces larger amounts of

active species, possibly by counter-balancing competitive coordination of the substrate to the metal precursor.

3.6. Temperature, Pressures

Considering the moderate activity of the Ru-based systems, experiments performed at higher temperatures with the $\text{RuCl}_2(\text{PPh}_3)_3/\text{Biphephos}$ system were first envisioned. The compositions profiles and selectivities obtained at 140 °C, instead of 120 °C, are presented in Figure 2 and Table 5. As expected, an increase of the global reaction rate was observed at 140 °C. However, the isomerization process was much more favored at this temperature, as compared to 120 °C; the amount of internal olefins increased from 22% to 40% and the selectivity in hydroformylation products concomitantly dropped from 76% down to 58%. The significantly larger amount of internal olefins that migrated at least twice (**1-int-x**; $x = 0/1^+ = 88:12$) underscores the importance of the isomerization process. The relative pseudo zeroth-order rates^[23] of hydroformylation vs. isomerization, as determined from the initial rates in Figure 2 and expressed as $k_{\text{HF}}/k_{\text{Iso}}$, decreased from 4.6 at 120 °C down to 1.4 at 140 °C; this corresponds to a difference in activation energies $E_{\text{a,Iso}} - E_{\text{a,HF}}$ of *ca.* 8 $\text{kJ}\cdot\text{mol}^{-1}$. It is noteworthy that even at the latter high temperature, conversion of the internal isomers **1-int-x** to aldehydes did not proceed at a noticeable rate (see Figure 2); this evidences the impossibility, at least with the present catalyst systems, to achieve a tandem isomerization-hydroformylation process [17]. Yet, the increase in temperature from 120 to 140 °C only had a minimal impact on the *l/b* ratio and, more surprisingly, on the chemoselectivity (in particular, larger amounts of alcohols may have been anticipated since this is a consecutive, more energy-demanding process).

Modification of the total and relative pressures is likely to influence the formation of active species and/or lead to different catalytic species, and eventually affect activities and selectivities [24]. Representative results of such variations in the $\text{RuCl}_2(\text{PPh}_3)_3/\text{Biphephos}$

system are presented in Table 6. When the total syngas pressure was varied in the range 10–40 bars, at a constant 1:1 CO/H₂ ratio, the reactions proceeded with just slightly decreased activity (from 5.0–6.1 min⁻¹ down to 3.5 min⁻¹) and quite similar selectivities (entries 25, 26 and 29). A much more significant decrease of the catalytic activity was observed at 90 bars, where the TOF dropped down to 0.6 min⁻¹; meanwhile, the amount of branched aldehyde was doubled while the chemoselectivity remained constant (entry 30). The same trends were observed upon changing the CO/H₂ ratio to 3:1 at a total pressure of 20 bars (entry 28). Obviously, excess CO is detrimental. On the other hand, when the CO/H₂ ratio was set at 1:3, the activity somehow decreased but most noticeably, the chemoselectivity for aldehydes dramatically decreased; in particular, larger amounts of alcohols **5** were formed, a non-expected result (entry 27).

3.6. Recycling

Attempts to recycle the catalyst and eventually improve on the catalytic productivity were conducted. We used the same procedure as the one positively evaluated for the analogous Rh-Biphephos system [15a]: the vacuum distillation of the crude reaction mixture can be readily achieved, allowing complete elimination of toluene solvent, and recovery of analytically pure aldehydes (along with residual internal olefins) and of a solid residue that contains the catalyst/ligand. To prevent deteriorated performance, the solid residue was concentrated and recovered under an inert atmosphere. Representative results obtained using this procedure are gathered in Table 7; the runs were conducted over long time period (124–190 h) to achieve high conversions. The results evidence that effective recycling can be achieved, maintaining a good chemo- and regioselectivity in favor of the linear aldehyde over at least three runs; the overall TON thus reached 55,000. The addition of a novel charge of fresh Biphephos in the recycling runs does not appear essential (compare run 2 and run 3). Yet, a ³¹P NMR monitoring of the reaction mixture was also performed, indicating that the Biphephos ligand slowly degrades under the reaction conditions (see the Supporting Information, Figure S1); similar observations were made with the Rh-Biphephos catalyst system [15a].

4. Conclusions

Combination of Ru(II) metallic precursors with Biphephos affords a selective catalytic system for the hydroformylation of 10-undecenitrile. The selectivity data: 75% of hydroformylation and *l/b* ratio up to 99:1, compare favorably with the very good performance of the rhodium- and iridium-based catalysts we previously reported [5c,15]. Formation of the hydrogenated olefin and of the alcohols resulting from the reduction of the desired aldehydes can be prevented by an adequate choice of the temperature and CO/H₂ pressures. These good chemo- and regio-selectivities are balanced with the lower activity (non-optimized hydroformylation TOF_{HF} of 2-7 min⁻¹), which are decreased by *ca.* 1-2 orders of magnitude (and not 5 orders as initially anticipated) in comparison with equivalent Rh-Biphephos systems (TOF_{HF} = *ca.* 80 min⁻¹), although the lower prices of ruthenium (*ca.* 15 times cheaper than Rh and Ir) must also be taken into account. This reduced activity did not hamper to achieve very high productivities (effective TurnOver Numbers for aldehydes, TON, up to 15,000 for batch experiments and up to 55,000 upon recycling). A limitation of these ruthenium systems is their incapacity, under the conditions suitable for selective hydroformylation, to promote isomerization of internal olefins, and hence to achieve a tandem isomerization-hydroformylation process as the Rh-Biphephos system is amenable to; we assume that this reflects the lower isomerizing ability of putative Ru-hydride species generated in these systems as compared to the corresponding Rh-hydride species in the Rh-based systems. Also, these ruthenium-based systems seem to be more versatile than the rhodium ones, as important isomerization or loss of regioselectivity can occur more easily if freshness of the catalyst precursors is not perfectly controlled.

Supporting Information Available

³¹P{¹H} NMR monitoring of reaction media.

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Figure caption

Figure 1. Diphosphine and diphosphite ligands used for ruthenium-catalyzed hydroformylation

Figure 2: Distribution of substrate and products (◆ **1**, ■ **1-int-x**, ▲ **2+3**) as a function of time at 120 °C and 140 °C in the hydroformylation of 10-undecenitrile by the $\text{RuCl}_2(\text{PPh}_3)_3/\text{Biphephos}$ system.

Scheme 1. Aldehydes and side-products arising from the hydroformylation-isomerization of 10-undecenitrile (**1**)

Table caption

Table 1. Variation of precursor in Ru-catalyzed hydroformylation of 10-undecenitrile.^a

Entry	Ru precursor	Time	1	1-int-x	1-int-x	2+3	2/3	4	5	Conv. 1	HF	TOF ^f
		[h]	[%] ^b	[%] ^b	$x=0/1^+$	[%] ^b	(<i>l/b</i>) ^c	[%] ^b	[%] ^b	[%] ^d	[%] ^e	[min ⁻¹]
1	$\text{RuCl}_2(\text{PPh}_3)_3$	48	10	22	98:2	63	99.1:0.9	4	0.3	89	76	6
2	$\text{RuCl}_2(p\text{-cymene})$	48	55	15	99:1	27	98.5:1.5	3	0.4	42	69	3
3	$[\text{Ru}(\text{COD})\text{Cl}_2]_n$	48	62	13	nd	23	99.1:0.9	2	traces	34	72	2
4	$\text{RuCl}_2(\text{DMSO})_4$	48	0	24	92:8	71	98.9:1.1	5	0.5	100	75	7
5	$\text{Ru}_3(\text{CO})_{12}$	64	45	16	99:1	36	99.2:0.8	2	0	51	75	3

^[a] Reaction conditions: $[\mathbf{1}]_0/[\mathbf{1-int-x}]_0$ (94:6) = 5.0 mmol, $[\mathbf{1}]_0/[\text{Ru}] = 20,000$, $[\text{Biphephos}]/[\text{Ru}] = 20$, toluene (5 mL), $\text{CO}/\text{H}_2 = 1:1$, $P_{\text{tot}} = 20$ bar, $T = 120$ °C. ^[b] Distribution (mol-%) of remaining **1**, internal alkenes **1-int-x** (residual or formed during the reaction; $x=0/1^+$ refers to the positioning of the internal C=C bond, $x = 0$ being 9-undecenitrile and $x = 1+$ referring to 8-, 7-, ...undecenitriles; please refer to Scheme 1), aldehydes **2** and **3**, hydrogenated product **4**, and alcohols **5** resulting from aldehydes reduction, as determined by ¹H NMR analyses.

^[c] Regioselectivity as determined by the linear-to-branched aldehyde ratio. ^[d] Conversion of **1** into **1-int-x** and **2-5**, calculated taking into account the quantity of **1-int-x** initially present in the substrate: $\text{Conv} = ([\mathbf{2}]_t + [\mathbf{3}]_t + [\mathbf{4}]_t + [\mathbf{5}]_t + [\mathbf{1-int-x}]_t - [\mathbf{1-int-x}]_0) / [\mathbf{1}]_0$. ^[e] Chemoselectivity as determined by the percentage of hydroformylation among all other competitive processes. ^[f] Overall TOF determined from the conversion of **1** over the whole reaction time.

Table 2: Solvent comparison in the Ru-catalyzed hydroformylation of 10-undecenitrile.^a

Entry	Ru precursor	solvent	time	1	1-int-x	1-int-x	2+3	2/3	4	5	Conv. 1	HF	TOF ^f
			[h]	[%] ^b	[%] ^b	$x=0/1^+$	[%] ^b	(<i>l/b</i>) ^c	[%] ^b	[%] ^b	[%] ^d	[%] ^e	[min ⁻¹]
6	RuCl ₂ (PPh ₃) ₃	toluene	48	10	22	98:2	63	99.1:0.9	4	0.3	89	76	6
7	RuCl ₂ (PPh ₃) ₃	CH ₃ CN	48	10	22	97:3	64	99.2:0.8	4	0.3	89	77	6
8	RuCl ₂ (PPh ₃) ₃	DMF	50	22	19	97:3	56	98.7:1.3	3	0	76	80	5
9	RuCl ₂ (PPh ₃) ₃	diglyme	44	41	20	99:1	37	98.8:1.2	2	0	55	72	4
10	RuCl ₂ (PPh ₃) ₃	DCE	48	77	14	96:4	4	98.0:2.0	4	traces	16	36	1
11	RuCl ₂ (DMSO) ₄	toluene	48	0	24	92:8	71	98.9:1.1	5	0.5	100	75	7
12	RuCl ₂ (DMSO) ₄	CH ₃ CN	48	1	25	93:7	70	99.1:0.9	4	0	99	76	7
13	RuCl ₂ (DMSO) ₄	DMF	47	5	31	94:6	60	98.6:1.4	4	0	95	68	7
14	RuCl ₂ (DMSO) ₄	diglyme	48	70	11	nd	16	97.5:2.5	3	nd	25	68	2

^[a] See Table 1.

Table 3: Ligand comparison in the Ru-catalyzed hydroformylation of 10-undecenitrile.^a

Entry	Ligand	Time	1	1-int-x	1-int-x	2+3	2/3	4	5	Conv. 1	HF	TOF ^f
		[h]	[%] ^b	[%] ^b	$x=0/1^+$	[%] ^b	(<i>l/b</i>) ^c	[%] ^b	[%] ^b	[%] ^d	[%] ^e	[min ⁻¹]
15	Biphephos	48	10	22	97:3	64	99.1:0.9	4	0.3	68	76	5
16	A4N3	66	30	20	97:3	49	98.7:1.3	1	traces	88	76	4
17	P(OPh) ₃	68	70	15	nd	15	81.9:18.1	1	traces	25	63	1

^[a] Metallic precursor: RuCl₂(PPh₃)₃ + 20 equiv. of ligand, except P(OPh)₃, 40 equiv.; otherwise, see Table 1.

Table 4: Influence of the ligand-to-metal ratio in the hydroformylation of 10-undecenitrile by the RuCl₂(PPh₃)₃/Biphephos system.^a

Entry	[L]/[Ru]	Time	1	1-int-x	1-int-x	2+3	2/3	4	5	Conv. 1	HF	TOF ^f
		[h]	[%] ^b	[%] ^b	$x=0/1^+$	[%] ^b	(<i>l/b</i>) ^c	[%] ^b	[%] ^b	[%] ^d	[%] ^e	[min ⁻¹]
18	0	88	92	7	99:1	0	-	0.4	0	1	0	-
19	2.4	90	59	12	97:3	27	99.1:0.9	2	0	37	79	1.4
20	6.0	53	49	17	99:1	32	98.8:1.2	2	0.3	47	71	2.9
21	10.5	73	40	22	97:3	35	99.1:0.9	3	0	57	66	2.6
22	20	48	10	22	97:3	64	99.1:0.9	4	0.3	88	76	6.1

^[a] See Table 1.

Table 5: Influence of temperature in the hydroformylation of 10-undecenitrile by the RuCl₂(PPh₃)₃/Biphephos system.^a

Entry	Temp	Time	1	1-int-x	1-int-x	2+3	2/3	4	5	Conv. 1	HF	TOF ^f
	[°C]	[h]	[%] ^b	[%] ^b	$x=0/1^+$	[%] ^b	(<i>l/b</i>) ^c	[%] ^b	[%] ^b	[%] ^d	[%] ^e	[min ⁻¹]
23	120	48	10	22	97:3	64	99.1:0.9	4	0.3	68	76	5
24	140	63	1	40	88:12	54	98.7:1.3	5	0.3	99	58	11 ^g

^[a] See Table 1 for experimental conditions except for T. ^[g] TOF value calculated at its maximal slope.

Table 6: Influence of total and relative pressures in the hydroformylation of 10-undecenitrile by the RuCl₂(PPh₃)₃/Biphephos system.^a

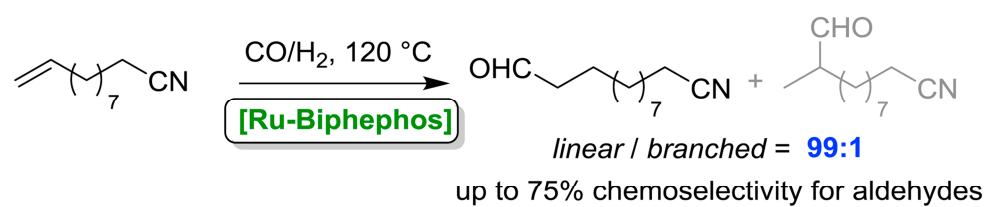
Entry	P _{tot}	CO/H ₂	Time	1	1-int-x	1-int-x	2+3	2/3	4	5	Conv. 1	HF	TOF ^e
	[bar]	ratio	[h]	[%] ^b	[%] ^b	$x=0/1^+$	[%] ^b	(<i>l/b</i>) ^c	[%] ^b	[%] ^b	[%] ^c	[%] ^d	(min ⁻¹)
25	10	1:1	63	24	23	97:3	50	99.3:0.7	3	1.1	74	72	3.9
26	20	1:1	48	10	22	97:3	64	99.1:0.9	4	0.3	89	76	6.1
27	20	1:3	55	58	18	99:1	14	98.9:1.1	5	5.7	38	40	2.3
28	20	3:1	65	71	13	99:1	13	98.8:1.2	1	1.6	24	60	1.2
29	40	1:1	46	43	15	98:2	39	98.9:1.1	2	0.6	54	77	3.9
30	90	1:1	71	81	8	98:2	10	97.9:2.1	1	0.0	13	78	0.6

^[a] See Table 1 except for the P value.

Table 7: Recycling of the RuCl₂(PPh₃)₃-Biphephos system over 3 runs in the hydroformylation of 10-undecenitrile.^a

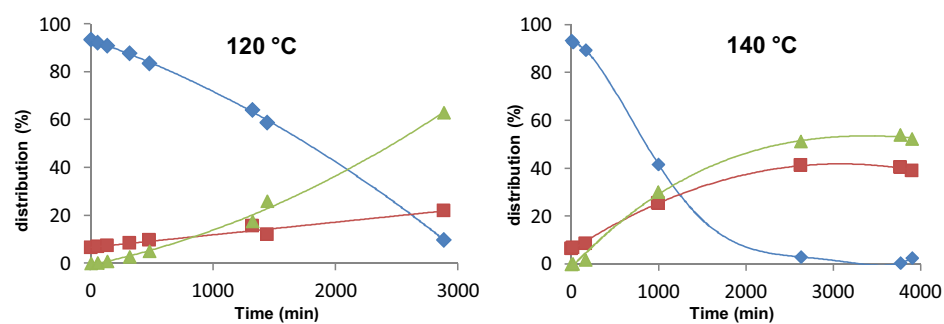
Run	Biphephos	time	1	1-int-x	2+3	2/3	4	5	Conv. 1	HF	TOF ^f
		[h]	[%] ^b	[%] ^b	[%] ^b	(<i>l/b</i>) ^c	[%] ^b	[%] ^b	[%] ^d	[%] ^e	[min ⁻¹]
1	-	190	14	31	45	98.8:1.2	5	5	84	57	1.6
2	+20 equiv ^g	150	6	28	62	99.3:0.7	3	1	94	71	2.2
3	-	124	3	27	64	98.9:1.1	5	1	99	72	2.7

^[a] See Table 1. ^[g] 20 equiv of Biphephos (vs. Ru) were added to the solid residue before starting the 2nd run.



TON up to 15,000 mol(aldehyde)·mol(Ru)⁻¹ and TOF up to 310 mol·mol(Ru)⁻¹·h⁻¹

fx1 .



gr2 .

*