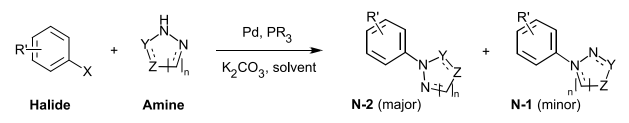


## Process Development Case Study WORK-UP

Design of a Washing Regime to Purge a Regioisomer

#### **INTRODUCTION**

An optimised palladium-catalysed **Buchwald-Hartwig amination** reaction was found to **reliably produce** the hetero-biaryl product with a regioisomeric ratio in solution of **92:8** in favour of **the desired N-2 isomer** at end of reaction (EOR).



**Crystallisation** using the prevailing conditions resulted in an **unacceptable level** of 1.5% of N-1 isomer still remaining in the isolated drug substance. Since the drug substance specification required <0.3% of N-1 isomer the challenge was to **improve the purity** further on work-up by **removing the undesired isomer** with **minimum loss** of the requisite N-2 **product**.

## **KEY OBSERVATION**

The aqueous wash after the cross-coupling reaction removed excess base and inorganic salts. Analysis showed that there had also been a **preferential partitioning of the unwanted N-1** isomer into the **aqueous wash**. A significant improvement in the isomer ratio to 95:5 for the organic phase was observed, with a concomitant ratio of 67:33 in the aqueous phase; **loss of product** to the aqueous phase was **7%** (**Table 1**).

	At EOR	After wash #1		After wash #2		After wash #3	
Temp. (°C)	Ratio*	Ratio*	% loss (total)†	Ratio*	% loss (total)†	Ratio*	% loss (total)†
25	92:8	95:5	18	97:3	59	99:1	61
80	92:8	95:5	7	97:3	12	99:1	18

\* ratio is of N-2 : N-1 isomers in organic phase; † % loss is cumulative from all sources

Table 1 - cumulative effects of aqueous washes at low and high temperatures

When **multiple water washes** were carried out, the ratio of N-2 to N-1 improved further to **99:1**. However, this also resulted in partial precipitation of the N-2 product at the interface, leading to **significant loss of product** from the organic phase and thus from the process. This was much **less pronounced at higher temperatures** due to increased solubility (18% loss over three washes at 80 °C versus 60% loss over three washes at 20 °C).

This observation showed that the first wash was more selective in extracting the undesired N-1 isomer and did not cause the precipitation of the N-2 product at the interface that was observed in following washes. These results were attributed to either the difference in the ionic strength and/or the pH of the initial aqueous wash that arose following the removal of the dissolved inorganic salts and excess base from the crude reaction mixture, respectively. To test this hypothesis further, the subsequent washes were designed to imitate the ionic and basic properties of the first wash.

### DISCUSSION

When **brine washes** were employed as substitutes for simple water washes, the loss of the **desired N-2 isomer** to the aqueous phase was **significantly reduced** over the course of three washes (11% compared to 18%) whilst the preference for extraction of the N-1 isomer over the N-2 isomer was retained (**Table 2**). In addition, **the partial precipitation of product** at the interface observed in the second and third water washes was **no longer observed** when using brine solutions. The individual washes (each 10 relative volumes) were designed with ionic strengths varying from 1-3% w/v sodium chloride to ensure maximum retention of the undesired N-1 isomer.

CatSci Ltd, CBTC2, Capital Business Park, Wentloog, Cardiff, CF3 2PX, United Kingdom



# Process Development Case Study WORK-UP

Design of a Washing Regime to Purge a Regioisomer

	At EOR	After w	vash #1	After wash #2		After wash #3	
Brine		1% w/v		3% w/v		3% w/v	
Phase	Ratio*	Ratio*	% loss (total)†	Ratio*	% loss (total)†	Ratio*	% loss (total)†
Organic	92:8	96:4	n/a	97:3	n/a	>98:2	n/a
Brine	n/a	55:45	neg.	65 : 35	3.1	90:10	11

\* ratio is of N-2 : N-1 isomers in either phase; † % loss is cumulative from all sources n/a = not applicable; neg. = negligible

Table 2 - cumulative effects of brine washes at 80 °C

To further evolve this concept, **alkaline brine solutions** were also investigated. A basic pH for the second and third washes was achieved by the dissolution of 1 equivalent of  $K_2CO_3$ . This was expected to reduce the loss of the desired N-2 isomer in the aqueous layer still further, and indeed this was found to be the case. Accordingly, **loss of the desired N-2 isomer** was limited to a total of only **5%** using such alkaline brine washes, whilst the selectivity for the N-1 isomer remained unaltered at >98:2 (**Table 3**).

	At EOR	After wash #1		After wash #2		After wash #3	
Alkaline brine		1% w/v		3% w/v plus K <sub>2</sub> CO <sub>3</sub>		3% w/v plus K <sub>2</sub> CO <sub>3</sub>	
Phase	Ratio*	Ratio*	% loss (total)†	Ratio*	% loss (total)†	Ratio*	% loss (total)†
Organic	92:8	95:5	n/a	97:3	n/a	>98:2	n/a
Brine	n/a	55:45	neg.	60:40	neg.	70:30	5

\* ratio is of N-2 : N-1 isomers in either phase;  $\uparrow$  % loss is cumulative from all sources n/a = not applicable; neg. = negligible

Table 3 - cumulative effects of alkaline brine washes at 80 °C

### **EXPERIMENTAL PROCEDURE**

All experiments were carried out in a nitrogen filled glove-box using nitrogen degassed solvents. The reactions were carried out on a scale of at least 2 g of starting Halide. The active catalyst was prepared out by heating the Pd pre-catalyst/ligand solution at 60 °C for 30 mins. This was transferred to the other reagents and the reactions were then stirred at 110 °C for 5 hours. The temperature of the reaction mixture (15 volumes) was reduced to 80 °C after completion of the reaction and the aqueous phase (10 volumes, water/brine/alkaline brine) was heated to 80 °C prior to addition. The resulting mixture was stirred for 20 minutes at 80 °C, then allowed to stand for 20 minutes to ensure full phase separation. Brine washes: wash 1 (1% w/v NaCl), washes 2 and 3 (3% w/v NaCl). Alkaline brine washes: wash 1 (1% w/v NaCl), washes 2 and 3 (3% w/v NaCl).

#### **SUMMARY**

**Optimisation of the work-up** successfully **improved the regioisomeric ratio** from 92:8 at EOR to >98:2 in solution with minimal loss of yield (~5%). The final crystallisation of the drug substance then resulted in <0.2% of the unwanted N-1 isomer, **within specification**. This was achieved through i) initial **experimental observation**; ii) **correct deduction** of its significance; iii) **quantitative analysis** of the organic and aqueous phases. Thorough understanding of the work-up process enabled successive improvements to be harvested. The **beneficial work-up** was fully **demonstrated** at CatSci (0.5 kg batch, 20 L scale) and subsequently at the customer's **manufacturing facility**.

Want to find out more? Contact us at **technical\_enquiries@catsci.com** or visit our website **catsci.com** 

CatSci Ltd, CBTC2, Capital Business Park, Wentloog, Cardiff, CF3 2PX, United Kingdom