

# SCAVENGING USING Silia*MetS®* AND Silia*Bond®* SCAVENGERS

Scavenging studies and results in one place



# Scavening of:

Antimony (Sb)
■ Cesium (Cs)
■ Copper ( <i>Cu</i> )
■ Iridium ( <i>Ir</i> )
■ Manganese (Mn)
<ul> <li>Osmium (Os)</li> </ul>
■ Palladium (Pd)
■ Platinum ( <i>Pt</i> )
■ Ruthenium ( <i>Ru</i> )
■ Tin ( <i>Sn</i> )
■ Uranium ( <i>U</i> )
<ul> <li>Multiple Metals</li></ul>
<ul> <li>Organic Impurities</li></ul>



Scavenging of Metal and Organic Impurities Studies







# Application Notes & Case Studies Silia*MetS* Metal Scavengers

Removal of Sb

# *In situ* speciation of dissolved inorganic antimony in surface waters & sediment porewaters

Antimony is a priority environmental contaminant, yet very little research on the behavior and fate of Sb in the environment has been reported. The primary source is mining and processing of Sb for industrial applications. To properly understand the processes controlling the movement of Sb through the environment, there is a need for tools to accurately measure the speciation of Sb, not just total concentration.

Both toxicity & mobility of Sb are affected by oxidation state, thus the accurate measurement of Sb speciation is essential for investigating the behavior of this contaminant in aquatic systems.

The authors present herein a diffusive gradients in thin films (*DGT*) technique, which utilizes a binding layer containing Silia*MetS* Thiol, for the selective measurement of Sb<sup>III</sup> in surface waters and sediment porewaters.

The mercapto-silica showed a high affinity for Sb<sup>III</sup>, with uptake efficiencies > 97 % and elution efficiencies of 86.9 +/- 2.6 %

The Silia*MetS* Thiol based DGT technique was also applied to measure porewater concentrations of Sb<sup>III</sup> and As<sup>III</sup> in a contaminated freshwater sediment at high resolution.

Initial and/or residual metal content was not disclosed.

Bennet, W. W. et al. Environ. Sci.: Processes Impacts, **2016**, *18*, 992-998 Griffith University, South East Queensland, Australia

#### Removal of Cs

# Amphoteric borylketenimines: versatile intermediates in the synthesis of borylated heterocycles

Reported herein was the 1<sup>st</sup> synthesis of amphoteric borylketenimines from ethynyl N-methyliminodiacetic acid (*MIDA*) boronate and sulfonyl azides via copper catalysis.

This work highlights the unexplored utility of borylketenimines in the synthesis of potentially bioactive molecules.

In situ Silia*MetS* Thiol (4 equiv) successfully removed the remaining Cs in deprotection reaction, which resulted in complete conversion and high selectivity.



Initial and/or residual metal content was not disclosed.

Yudin, A. K. *et al. Chem. Eur. J.*, **2017**, *23*, 9711-9715 University of Toronto, Ontario, Canada





# Silia*MetS* Imidazole in the synthesis of polymer films for measurement of optical properties

An asymmetric bifunctional monomer having both an unprotected alkynyl group and a vinyl ether (*VE*) polymerization of VEEP smoothly proceeded in a controlled manner under a living cationic polymerization condition to give alkyne-substituted polyVE (*poly-VEEP*) without any protection of the pendant alkynyl function.

In the click reaction of precursor macromonomer MA-PVEEP, the resultant polymer MA-P(*VE-Mal*) was purified from contaminating copper with Silia*MetS* Imidazole (0.192 mmol, 1.13 equiv).



### Removal of Cu

### Synthesis of a multivalent glycoligand against avian and human influenza viruses

A glycopolymer bearing  $\alpha$ 2,3-linked sialyltrisaccharides was synthesized by living radical polymerization using a glycomonomer prepared by a protecting-group-free process, direct azidation of the free sialyllactose, and subsequent azide-alkyne cycloaddition.

The prepared glycopolymer with pendant 3'-sialyllactose moieties strongly interacted with both avian & human influenza viruses.

SiliaMetS Imidazole (5 equiv for Cu) was successful in removing contaminating Cu, with only 5 equiv, O.N. at r.t.





Removal of Ir

# Hydrogen-borrowing and interrupted-hydrogen-borrowing reactions of ketones and methanol catalyzed by iridium

The use of catalytic  $[{Ir(cod)Cl}_2]$  was shown to facilitate hydrogen-borrowing reactions of ketone enolates with methanol at 65°C. An oxygen atmosphere accelerated the process, and when combined to a bulky monodentatephosphine ligand, the catalytic cycle could be interrupted by preventing the enone reduction.

Subsequent addition of pronucleophiles to the reaction mixture allowed a one-pot methylenation/conjugate addition protocol to be developed: the enone and methoxy adduct were not isolated but reacted *in situ* with an external nucleophile and extra base. In most cases they found to be beneficial to treat the crude reaction mixture with the Ir scavenger Silia*MetS* DMT while stirring the solution, open to the atmosphere, for 1 hour, before the addition of base and nucleophile. The team suggested that the removal of most of the metal catalyst from solution and prevented complications caused by methanol oxidation during the second phase of the reaction.



Initial and/or residual metal content was not disclosed.

Donohoe, J. et al. Angew. Chem. Int. Ed., 2014, 126, 1-5
 Donohoe, J et al., Angew. Chem. Int. Ed., 2015, 54, 1642-1645
 University of Oxford, Oxford, United Kingdom
 Novartis Institute for Biomedical Research, Cambridge, United States

Removal of Mn

# Diastereoselective synthesis of cis-N-Boc-4-aminocyclohexanol with reductive ring opening method using continuous flow

N-protected cis-4-aminocyclohexanol derivatives are valuable intermediates in the syntheses of APIs.

Compared with previously reported batch processes, the present multistep flow process is much more efficient in synthetizing valuable 4-aminocyclohexanol derivatives, and avoids some isolation and purification steps.

In this procedure, a nitroso hetero-Diels-Alder cycloaddition step using  $MnO_2$  as the oxidant was fully integrated to a novel continuous flow process for the hydrogenation and ring opening of cycloadducts to the corresponding cis-4-aminocyclohexanols using H-Cube Pro.



It was shown that Raney Nickel was a significantly more performant catalyst compared to 10 % Pd/C and 5 % Ru/C in this last cleavage, and highly selective ( $\geq$  99 % toward final cis-4-aminocyclohexanol). Yet, the Raney Nickel cartridge lost its activity after a few measurements. A possible manganese leaching into the reaction mixture was presumed as the cause for catalyst deactivation. Based on the atom absorption measurements the average extent of manganese was 15 ppm in the reaction mixture. Filling the end of the MnO<sub>2</sub> column with Silia*MetS* AMPA as a Mn scavenger allowed the Raney nickel catalyst activity to be kept constant for longer time.

Initial and/or residual metal content was not disclosed.

Szabó, B. *et al. J. Flow Chem.*, **2019**, *9*, 13-17 University of Technology and Economics, Budapest, Hungary





### Osmium scavenging with SiliaMetS

Osmium tetroxide  $(OsO_4)$ , is a very reliable and powerful reagent for the cis-dihydroxylation of alkenes. However, osmium compounds, in particular  $OsO_4$ , are highly poisonous, even at low exposure levels, and must be handled with appropriate precautions.

Therefore, it is important to efficiently remove residual osmium from products of interest. A scavenging study on three organic reactions involving osmium reactants were performed. The metal scavenging efficiency of Silia*MetS* is highlighted in the following table.

$C_6H_{13}$ $\longrightarrow$ $H_{13}C_6$	OH OH H <sub>13</sub> C <sub>6</sub>	AD-Mix-β	Н <sub>13</sub> С <sub>6</sub> ОН ОН	H <sub>13</sub> C <sub>6</sub> OSO NaIC	H <sub>13</sub> C <sub>6</sub>	
Dihydroxylation Reaction [OsO <sub>4</sub> ] [i		Sharpless Dihydrox Potassium osmate (K <sub>2</sub>	ylation Ds <i>O</i> 2(OH)4]	Lemieux-Johnson Oxidation $[NalO_{a}, OsO_{a}]$		
Y.	Osmium Scavenging using Silia <i>MetS (in %</i> ) at 22°C					
SiliaMotS	Dihydroxylation	Sharpless Di	nydroxylation	Lemieux-Johnson Oxidation		
SilidivietS	4 equiv, 4 h	8 equiv, 4 h	8 equiv, 16 h	8 equiv, 4 h	8 equiv, 16 h	
Silia <i>MetS</i> Thiol	87	> 98	> 98	87	92	
SiliaMetS Cysteine	89	> 98	> 98	87	91	
Silia <i>MetS</i> DMT	92	97	> 98	87	91	
SiliaMetS Imidazole	87	> 98	> 98	89	91	
Initial Os Concentration ( <i>in EtOAc</i> ) 132 ppm 25 ppm			ppm	21 г	opm	

Note: > 98 % of scavenging means < 0.5 ppm of osmium.

All scavengers were equally effective for Sharpless dihydroxylation or Lemieux-Johnson oxidation. As for simple dihydroxylation, Silia*MetS* DMT was slightly more efficient.



# Identification of a potent, highly selective, and brain penetrant phosphodiesterase 2A inhibitor clinical candidate

Computational modeling was used to direct the synthesis of analogs of previously reported phosphodiesterase 2A (*PDE2A*) inhibitor with an imidazotriazine core to yield compounds of significantly enhanced potency.

The analog PF-05180999 was subsequently identified as a preclinical candidate targeting cognitive impairment associated with schizophrenia.



Silia*MetS* Thiol was used to remove Pd from the palladium-catalyzed C–H arylation on the pyrazole C-5 position with heteroaryl halide to synthesize the PF-05180999. It was found that C–H arylation chemistry was superior to Suzuki coupling for heteroaromatic derivatives, likely due to instability of the required heteroaryl boronates.

Initial and/or residual metal content was not disclosed.

⊟ Helal, C. et al. J. Med. Chem., 2018, 61, 1001-1018 Research API, Pfizer Global Research & Development, Connecticut, United States



#### Removal of Pd

# N-annulated perylene diimide dimers: controlling structural conformation & impact on physical, electronic, optical & photovoltaic properties

The geometry of organic  $\pi$ -conjugated small molecules can impact the morphology of blended-thin films and subsequent performance in opto-electronic devices.

The role of molecular conformation of perylene diimide (*PDI*) dimers designed to act as non-fullerene acceptors in organic solar cells was studied.

SiliaMetS DMT removed all traces of Pd to generate Sonogashira product in near quantitative yield.

This was mandatory in the synthesis as it was found that the subsequent deprotection to form PDI-Ac was found to be sensitive to the amount of Pd remaining in the sample.



### Removal of Pd

# Suzuki-Miyaura cross-coupling synthesis of O6-benzyl glaziovianin A

 $O^6$ -benzyl glaziovianin A is a potent cytotoxic substance & a potent inhibitor of  $\alpha$ , $\beta$ -tubulin polymerization. Following a Suzuki-Miyaura cross-coupling, Pd-containing  $O^6$ -benzyl glaziovianin A was purified using Silia*MetS* Thiourea.

Purification was done stirring the API with SiliaMetS Thiourea at r.t. in CHCl<sub>3</sub> for 12 h.

Less than 1 equiv. of SiliaMetS (0.3 equiv) was needed for scavenging!



Hayakawa, I. *et al. Bioorg. Med. Chem.*, **2016**, *24*, 5639 Okayama University, Okayama, Japan University of Tsukuba, Tsukuba, Japan





# Synthesis of Mavatrep: a potent antagonist of transient receptor potential vanilloid-1

Mavatrep, a potent transient receptor potential vanilloid-1 (*TRPV1*) antagonist, was prepared in four steps in 63% overall yield. One of the two key synthetic transformations was the last step Suzuki coupling of an imidazole with an oxaborolole moiety.

Repeated attempts to carry out the Suzuki coupling reaction under the original conditions in presence of  $Pd(dppf)Cl_2$  as a catalyst resulted in very poor yields (5-35 %) and formation of high levels of impurities requiring tedious purification. This poor outcome was even worse on larger scale.

After screening various experimental conditions, the best result was obtained by the use of Pd(dtbpf)Cl<sub>2</sub> in DME in the presence of aqueous Na<sub>2</sub>CO<sub>3</sub> which led to a remarkable improvement in both reaction rate and yield. The crude salt was obtained in 80-90% as a brownish solid that typically contained about 1400-1600 ppm of residual palladium. Out of the six screened Silia*MetS* (*Diamine, TAAcOH, TAAcOH, Thiol, Thiourea and Triamine*), Thiourea proved to be the most effective palladium scavenger.

(1.05 equiv)



Pd(dppf)Cl<sub>2</sub> (0.03 equiv) DME / aq. Na<sub>2</sub>CO<sub>3</sub> 5 h , 80°C 2. Silia*MetS* Thiol (30 equiv) Silia*MetS* Thiourea (40 equiv) MeOH/EtOAc, 54-56°C, 1.5 h

ÒН

84 % Pd content: < 10 ppm

However, an unexpected result was found by the synergistic effect of a combination of two Silia*MetS*, Thiol and Thiourea when used in concert, which was not apparent from the screening array. Their use in combination left the API with less than 10 ppm of Pd.

Wells, K. M. et al. Org. Process Res. Dev., 2015, 19, 1774-1783 Janssen Research and Development, Pennsylvania, United States







Removal of Pd

# A Pfizer case study: palladium removal using Silia*MetS* Thiol after a Buchwald-Hartwig amination

Silia*MetS* Thiol was employed by researchers at Pfizer for Pd removal during the preparation of a naphthalenopiperazine HCl salt. The product from the Buchwald-Hartwig amination of naphthyl bromide with Boc-piperazine was telescoped as a toluene solution and the Boc protecting group was subsequently cleaved with HCl gas to afford the HCl salt containing over 1,300 ppm Pd.

A water / THF solution of this material was then treated with Silia*MetS* Thiol (50 *wt* %) at 35°C for 17 h and, following crystallization from water / THF.



### Conclusion

The HCl salt was obtained with a 90 % yield with only 2 ppm Pd. This chemistry was demonstrated on a kilogram scale.

■ J. Magano *et al.*, *J. Synth. Commun.*, **2008**, *38*, 3631-3639 Research API, Pfizer Global Research and Development, Connecticut, United States Separation Sciences, Pfizer Global Research and Development, Connecticut, United States Research API, Pfizer Global Research and Development, Michigan, United States Supply Chain API, Pfizer Global Research and Development, Michigan, United States





# Circumventing an unexpected catalyst poisoning in a Suzuki coupling

In one project, ring methoxylation of a chloronicotinate was achieved by scientists at Dart NeuroScience and Accela ChemBio, followed by a one-pot Suzuki-coupling to yield a biaryl intermediate, one that was needed on a KG scale in further synthetic steps.



Consistent and excellent results were first obtained from R&D (*mg-scale*) up to a first cGMP process-scale tryout (6.4 kg). Yet, subsequent couplings on other batches gave practically no expected birayl compound. Instead, the first methoxylation product simply hydrolyzed from its ester form to the corresponding carboxylic acid!



After investigation, the only time the team could reproduce the non-reactive behavior of the reaction was upon intentionally omitting the  $Pd(OAc)_2$  catalyst – hence the idea of catalyst poisoning – but the reaction could still be conducted by adding some more palladium in the reaction mixture (*up to 5 %*). This also leads to downstream issues, as higher palladium loading also implies higher residual palladium, which can greatly impair subsequent reactions.

Among all metal removal strategies, it was the use of Silia*MetS* Thiol that proved to be the best protocol for a simple, fast and efficient removal of palladium. The Pd levels were thus lowered below 200 ppm. The team concluded that "*The subsequent Pd removal step was carried out as planned, and the remainder of the GMP synthesis proceeded well to provide final API in the desired quantity and purity*", as the expected biaryl intermediate was isolated in 73 % yield and 98 % purity.

Initial and/or residual metal content was not disclosed.

Liu, J. et al. Org. Process Res. Dev., **2018**, 22, 111-116 Dart NeuroSciemce L.L.C., California, United States Accela ChemBio Ltd, Shanghai, China

#### Removal of Pd

# Development and scale-up to improve an API's physiochemical and bulk powder properties

Purification process for TAK-117, a selective PI3Ka inhibitor currently in Phase 1b of clinical trials, was developed and greatly improved the overall purity, recovery, physiochemical and bulk powder properties of the isolated product, which was obtained after a Suzuki coupling.



A purification process is required to control the Pd levels within International Conference on Harmonisation (ICH) guidelines.

The revised process was found to produce purified material across a total of 6 batches at 15 and 30 kg scales. Original protocol subjected TAK-117 to 2 treatments with decreasing quantities of GCB and Silia*MetS* Thiol scavenger (20 wt %) each then 10 wt % each). New protocol uses Silia*MetS* Diamine (80 wt %) and GCB (20 wt %).

Initial and/or residual metal content was not disclosed.

Papageorgiou, C. D. *et al. Org. Process Res. Dev.*, **2018**, *22*, 296-305 Takeda Pharmaceuticals International Co., Process Chemistry, Cambridge, United States



Removal of Pd

# Discovery of small molecule splicing modulators of survival motor neuron-2 (*SMN2*) for the treatment of Spinal Muscular Atrophy (*SMA*)

Spinal muscular atrophy (*SMA*), a rare neuromuscular disorder, is the leading genetic cause of death in infants and toddlers. SMA is caused by the deletion or a loss of function mutation of the survival motor neuron 1 (*SMN1*) gene. SMN2 (*survival of motor neuron 2*) is a closely related gene which can partially compensate for the loss of SMN1, however coding for a less stable SMN protein.

In recent years, modulating SMN2 pre-mRNA splicing has greatly improved SMA treatments by means of the discovery of LMI070 / Branaplam, a small molecule stabilizing interaction between the spliceosome and SMN2 pre-mRNA.

An efficient two-step synthesis of Branaplam (*from an advanced intermediate S2*) was developed to avoid late stage demethylation which proved challenging upon scaling. Suzuki coupling occurred between pinacol ester of (4-chloro-2-hydroxyphenyl)boronic acid and pyridazine intermediate S2 in the presence of PdCl<sub>2</sub>(dppf), CH<sub>2</sub>Cl<sub>2</sub> and sodium carbonate, followed by a second Suzuki coupling catalyzed by BrettPhos palladacycle with 1-Boc-pyrazole pinacol boronate to form the Boc protected intermediate. After stirring overnight in the presence of 1M HCl, and then treated with Silia*MetS* DMT (*5 mmol, 2 equiv*) for 20 h, Branaplam was obtained in good overall yield (*65 %*).



Cheung, A. K. et al. J. Med. Chem., 2018, 61, 11021-11036 Novartis Institutes for Biomedical Research & Novartis Pharmaceuticals, Cambridge, United States

Removal of Pd

# Discovery and synthesis of a direct activator of AMP-Kinase PF-06409577 against diabetic nephropathy

Adenosine monophosphate-activated protein kinase (*AMPK*) is a protein kinase involved in maintaining energy homeostasis within cells. PF-06409577 has been discovered as an indole acid with excellent balance of properties and has now been advanced to first-in-human trials for the treatment of diabetic nephropathy. Key indazoles in the synthesis were prepared by Suzuki coupling to some commercially available 5-bromoindazoles.

Silia*MetS* Thiol was used to remove remaining Pd after only 30 min of stirring.



Initial and/or residual metal content was not disclosed.

Cameron, K. et al. J. Med. Chem., **2016**, 59, 8068-8081 Pfizer Worldwide Research & Development, Massachusetts & Connecticut, United States





# Development of a scalable synthesis of an IDO<sup>1</sup> inhibitor featuring a stereospecific Suzuki coupling of a cyclopropane carboxylic acid

A modified synthetic route to an IDO inhibitor was developed, which synthetic steps comprised a chemoselective nitro reduction, a Miyaura coupling (*Scheme 1*) followed by a key stereospecific Suzuki coupling (*Scheme 2*). A systematic evaluation of the Suzuki conditions led to the identification of a robust catalyst/ligand/base combination to reproducibly duplicate the coupling reaction on large scale, without recourse to chromatography given optimized purification methodology.

Post-reaction treatment included activated charcoal (*for pigments, inorganics...*) and Silia*MetS* Thiol (*for residual Pd*): following the reaction work-up, the organic layer was stirred over activated charcoal (*125 g*) at 20 - 30°C for 2 h. Silia*MetS* Thiol (*1.3 kg*) was charged to the mixture, and the slurry was stirred for 16 h at ambient temperature and filtered through Celite.



Scheme 2

Purification of final IDO inhibitor was done by treatment with activated charcoal and Silia*MetS* Thiol for removal of pigments, inorganics and residual palladium. After the work-up, the organic layer was distilled under reduced pressure, and the resultant brown viscous liquid was dissolved in MTBE (8 *L*).

Activated charcoal (40 g), silica gel (300 g), and Silia*MetS* Thiol (300 g) were added into this solution and stirred at 20 - 30°C for 18 h. The slurry was filtered through Celite.

Overall, the API was delivered in high quality (> 99 %) and good yield (42 %, less than 10 ppm of Pd) for a 5-step synthesis.

<sup>1</sup> Indoleamine-2,3-dioxygenase is a heme-containing intracellular enzyme

Vaidyanathan, R. et al. Org. Process Res. Dev., **2018**, 22, 888-897 Chemical Development and API Supply, Bristol Myers Squibb Research and Development Center, India



#### Removal of Pd

# Improved synthetic route and scale-up of GSK973 using Silia*MetS* Thiol for Pd removal

The team at GSK worked on improving the synthetic route to GSK973 for larger scale synthesis in order to provide sufficient material for pre-clinical efficacy and safety studies for oncology trials. GSK973 is an inhibitor of the second bromodomains.

Overall, this study managed to:

- Reduce the number of steps (from 10 to 7)
- Transform the synthesis into a more environmentally friendly, greener process due to solvent changes and total quantities
- Improve the synthesis' reliability and scalability
- Increase overall yield (from 10 to 25 %)
- Lower the dependency for flash chromatography at large scales

One of the objectives was to reduce the number of steps. The carbonylation/saponification/amide coupling sequence was replaced with a one-step aminocarbonylation. Basing themselves on a set of conditions previously optimized for an aminocarbonylation of another compound of interest, the researchers reused and adjusted the parameters for the synthesis of GSK973 on a 300 mg scale prior to scaling up. The adjustments were done in accordance with GSK's guidelines on green chemistry. Good yields were obtained at the small scale.

While scaling-up, lower conversions arose due to the limited solubility of the compound in the solvent used (*n-butyl* acetate). Switching the solvent to 1,4-Dioxane and increasing the catalyst loading, succeeded in converting 100 % of the starting material, with 87 % isolated yield on the 5 g scale and 95 % isolated yield on the 45 g scale.



Reagents and conditions: (a) CO (1 atm), 5 mol %  $(Pd(OAc)_2, 5 mol \% Xantos, 1.3 equiv amine HCI, 2.5 equiv 2,6-lutidine, 1,4-dioxane, 85°C, 16 h, 95 % (97 % pure by LCMS); (b) Si thiol resin, CH<sub>2</sub>Cl<sub>2</sub>/MeOH, room temperature, then trituration in EtOAc, 83 %.$ 

While the product obtained at this step was initially purified by flash chromatography, at a larger scale, flash chromatography could be avoided by scavenging the residual Pd catalyst using Silia*MetS* Thiol. The recovered product was simply triturated, filtered, and concentrated under vacuum to give the resulting product with a yield of 83 % with over 99 % of purity. Both enantiomers (± *in Figure*) were finally separated by chiral preparative HPLC, and enough material was recovered for the subsequent in vivo efficacy and toxicology studies.

Alder, C. et al. Org. Process Res. Dev., 2022, 26, 365-379





# Pd scavenging after a Suzuki-Miyaura coupling using a GlaxoSmithKline published reaction

A metal scavenging study was performed following the synthesis of a key synthetic intermediate obtained by the Suzuki-Miyaura coupling presented in scheme below. Various parameters were investigated including the efficiency of Silia*MetS* in different formats, scavenging kinetics, intermediate recovery and purity.



### Small-Scale Scavenging (Synthesis Scale ~ 5 g)

The table below shows the most efficient Silia*MetS* Metal Scavenger products for the treatment of the reaction mixture after work-up in both bulk and fixed-bed mode bed (*pre-packed SPE cartridges*).

	SiliaMetS Scavenging Efficiency & Intermediate Recovery Results (in %)					
Scavengers	Batch Reactor Mode (Bulk)5 equiv, 4 h, 22°C5 equiv, 4 h, 40°C		Fixed-Bed Mode (SPE) 6 mL / 1 g	Intermediate Recovery		
Silia <i>MetS</i> Thiol	95	> 99	98	> 99		
SiliaMetS Thiourea	83	93	99	98		
SiliaMetS Cysteine	84	91	97	> 99		
Silia <i>MetS</i> DMT	97	> 99	> 99	98		
Initial Pd Concentration	179 ppm in MTBE		76 ppm in Toluene	-		

#### Scavenging Conclusion

Addition of 5 equivalents of Silia*MetS* products for 4 h at the end of the reaction reduces the residual metal concentration to single digit ppm.

#### Recovery & Purity Conclusion

Palladium was completely removed, while the organic compound was not sequestrated by Silia*MetS* products. No impurities were released.

### Larger Scale Scavenging (Synthesis Scale ~ 55 g)

Silia*MetS* Metal Scavengers in pre-packed Silia*Sep* Flash Cartridges are a great alternative for metal removal at process development scale. These cartridges offer excellent scavenging efficiency as shown by the results in associated table.



After the first run, almost all the palladium was captured.

After three runs, less than 1 ppm remained in solution.

#### <u>Results</u>

SiliaSep Scavenging Results (in %)				
Run # Scavenging				
1	97			
2	99			
3	> 99			

Experimental	<b>Conditions</b>
-	

Scavenger Used	Silia <i>MetS</i> Thiol		
Cartridge Size	120 g		
Nb. Equivalent	25 equiv		
Solution Volume	1 L		
Flow Rate	40 mL/min		

Initial Pd Concentration: 700 ppm in AcOEt

J. F. Toczko et al., Org. Proc. Res. Dev., 2008, 12, 896-899 Chemical Development, GlaxoSmithKline, North Carolina, United States





# Multi-kilo delivery of AMG 925 featuring a Buchwald-Hartwig amination and processing with insoluble synthetic intermediates

Amgen describes the development of a synthetic route to manufacture the drug candidate AMG 925 on kilogram scale. The synthetic route features a Buchwald-Hartwig amination using BrettPhos as ligand and conducted to afford 12 kg of product in a single batch.

Yet, the target levels of heavy metal in the drug substance AMG 925 were < 100 ppm, much lower than that found in the crude Buchwald-Hartwig product (700 - 800 ppm).



After the screening of potential scavengers, Silia*MetS* Thiourea was selected as the scavenger of choice since it was the most effective. At 100 wt % relative to crude product, in 80 % (v/v) toluene/MeOH and a total of 15 volumes of solvent, it afforded product in 90 % with less than 2 ppm of residual Pd.

Caille, S. *et al. Org. Process Res. Dev.*, **2015**, *19*, 476-485 Amgen Inc., Thousands Oaks, California

Removal of Pd

# Suzuki-Miyaura cross-coupling synthesis of GDC-0084, a glioblastoma multiforme cancer inhibitor

GDC-0084 is a BBB (*Blood-Brain Barrier*)-inhibitor of the PI3K pathway, where GBM cancer cells reside.

In order to remove residual palladium from the API, a scavenging study using 40 functionalized resins, silicas & activated carbon was performed. Representative samples containing 2,400 ppm Pd were exposed to Silia*MetS* Scavengers (*20 wt % loading, 14 h*) in a solvent mixture of THF and HOAc (67:33).

Both SiliaMetS Thiol & Thiourea were chosen for lowering residual Pd to ≤16 ppm in G-scale experiments.

On KG-scale, a mixture of the two scavengers was used.





Removal of Pd

# Scalable Zn-catalyzed regioselective allylation of 1 methyl-1- $\alpha$ -rhamnopyranoside

The construction of C–N bonds by Pd-catalyzed cross coupling of amines with aryl halides, pseudohalides, and aryl ethers is an uncontested tool for the synthesis of functionalized diaryl amines (*widely found in pharmaceuticals, natural products and so on*).



Zinc salts have been shown to promote the Buchwald-Hartwig coupling of azaindoles and azaindazoles with heteroaryl chlorides to provide the corresponding 1-aryl-1H-azaindoles and 1-aryl-1H-azaindazoles.

The optimal Zinc salt was shown to be  $Zn(OAc)_2$ , because of high conversion, availability, low molecular weight and cost. Silia*MetS* Thiol successfully removed traces of palladium.

Initial and/or residual metal content was not disclosed.

Vaidyanathan, R. et al. J. Org. Chem., 2017, 82, 7420-7427 Chemical Development and API Supply, Bristol Myers Squibb Research and Development Center, India

#### Removal of Pd

### Palladium-mediated synthesis of a near-infrared fluorescent K+ sensor

Potassium (K+) exits electrically excitable cells during normal and pathophysiological activity.

Currently, K+-sensitive electrodes and electrical measurements are the primary tools to detect K+ fluxes. A near-IR, oxazine fluorescent K+ sensor (*KNIR-1*) with a dissociation constant suited for detecting changes in intracellular and extracellular K+ concentrations were synthetized, via a Sonogashira coupling between a halogenated triazacryptand (*TAC*) and a tetrahydroquinoline. The reaction proceeded smoothly.

Remaining palladium in the resultant alkyne was removed using Silia*MetS* DMT, as failure to remove all of the palladium impurities in the following reduction reaction resulted in cleavage of the tetrahydroquinoline from the TAC K+ binding domain during reduction.



& Assumption College, Massachusetts, United States





# Synthesis of EDOT-containing polythiophenes and their properties in relation to the composition ratio of EDOT

Polythiophenes composed of 3,4-ethylenedioxythiophene (*EDOT*) and 3-hexylthiophene (*3HT*) with different composition ratios of EDOT in the repeat unit of the polymer backbone were synthesized by polycondensation reactions. The optical, electrochemical and charge transport properties of the polymers were compared and found to be well correlated with the EDOT composition ratio.

Three kinds of polythiophenes containing 3HT and different EDOT composition ratios were synthesized by polycondensation reactions using direct C–H arylation and Stille coupling, from which Pd residues was removed with Silia*MetS* DMT.



■ Imae, I. *et al. RSC Adv.*, **2015**, 5, 84694-84702 Hiroshima University, Hiroshima, Japan





# A Tetraphase case study: palladium scavenging in the development of the first fully synthetic fluorocycline using Silia*MetS* DMT

Process research and development of the first fully synthetic broad spectrum fluorotetracycline in clinical development was reported by Tetraphase Pharmaceuticals. The key reaction was a Dieckmann condensation between a suitable substituted aromatic moiety and a cyclohexanone derivative. Subsequent hydrogenolysis was extensively studied, using a Pd/C catalyst. Without any treatment, residual palladium levels as high as 2,000 ppm were detected.

Silia*MetS* DMT was found to be an excellent metal scavenger to reduce the residual Pd content to more than acceptable levels.



	Scale-Up of Hydrogenation and Pd Scavenging Results							
Entry <sup>1</sup>	Hydrogenation Time (h)	Degenation Time (h)Time for Slurry in EtOH / $H_2O(h)$ Yield (%)		Pd Content (ppm) (after treatment)				
1	12	2	82	0.4				
2	4	17	73	2				
3	7	2	77	0.39				
4	10	2.5	79	1.11				
5	11	4	85	< 0.2				

1 6.2 to 10 wt % Pd/C was used

### Conclusion

The reaction was run in THF / MeOH (3.3/10 v/v) under nitrogen with 10 wt % Pd/C. Without any treatment, residual palladium levels were as high as 2,000 ppm, but after stirring with 50 or 100 wt % Silia*MetS* DMT in MeOH for 2 - 3 h, residual levels were consistently below 1.5 ppm.

M. Ronn et al., Org. Proc. Res. Dev., **2013**, 17, 838-845 Tetraphase Pharmaceuticals Inc., Massachusetts, United States







### An Amgen case study: Palladium removal using various resins

Amgen published a chapter in "Catalysis of Organic Reactions" related to the use of scavengers for the removal of palladium in small to multi-kilogram production scales.

In this study, several parameters were evaluated, such as scavenging efficiency, influence of the scavenger loading and loss of product to adsorption (*recovery*). The study was based on a palladium-catalyzed sulfonamide coupling and scavenger screening was performed at both room temperature and 65°C using 31 different scavengers.



### Amgen Scavenger Screening Results

**Conditions**: 20 mg of each scavenger (20 % w/w) were placed in 2 mL HPLC vials each containing 1 mL of crude reaction mixture containing 100 mg of product. Each vial was sealed and agitated overnight. Initial palladium concentration was 423 ppm.

The best scavenger identified during their study was the Silia*MetS* Thiourea providing the lowest Pd content (*residual palladium concentration: 3 % or < 14 ppm*) without product sequestration. They mentioned that Silia*MetS* Thiourea was used extensively in early process development work.



### Cost Comparison for Most Efficient Scavengers (≥ 80 %)

At pilot-plant scale, the optimal compromise between the cost per ppm removed and the scavenging efficiency is crucial.

Results highlighted by the graph reduced the number of options to only four candidates for further evaluation: in pole position Silia*MetS* Thiourea, followed by TMT, TMT-Na<sub>3</sub> and Smopex 234.



Note: SiliaMetS DMT does not appear in this study because it had not been commercialized yet at the time of the study.



### **Top Four Scavengers Overview**

A screening validation was conducted on 1-g scale purification (*10 mL of solution*) with 20 % w/w of the four best scavengers at 65°C overnight. After filtration, residual metal concentration was analyzed by ICP-MS and product recovery was determined by HPLC. Silia*MetS* Thiourea was chosen for the large scale purification.

Screening Validation Results on Top 4 Scavengers							
Scavengers	Residual Metal Concentration ( <i>ppm</i> )			Product	Commentary from Amgen		
	Solution	in Solution	in Solid Product	Recovery			
SiliCycle Thiourea	14	11	158	102 %	Best performance		
ТМТ	33	15	264	104 %	Fine in suspension, filterability concerns on scale		
Smopex 234	36	38	496	84 %	Favorable cost but product recovery inadequate		
TMT-Na3	85	81	1 555	78 %	Very basic compounds ( <i>not effective with base-sensitive groups</i> ) and low recovery		
Initial Concentration	423 ppm	381 ppm	3,577 ppm	-	-		
Purification Scale	100 mg	1 g	1 g	1 g	-		

Please see Amgen's chapter for further details (see reference below).

#### Amgen's Conclusion

"Scavengers offer a practical and expedient option for removal of palladium from process streams to ensure quality of organic products... The screening protocol involves treatment of a candidate process stream with 20 % w/w scavenger on product at both room temperature and 65°C followed by analysis of Pd and product adsorption. High-temperature treatment increased the efficiency of Pd removal... Evaluation of process costs is key to identifying Pd removal solutions. Whiles cavengers add cost to a process, their use is often justified by the speed to production in early phase development."

J. Allgeier et al., « Application of Scavengers for the Removal of Palladiumin Small Lot Manufacturing », Catalysis of Organic Reactions, Chapter 5., Amgen Inc., Thousand Oaks, California





### The effect of variation of phosphorous ligand nature on scavenging

Even for the same metal, a variation in the scavenging efficiency can be observed depending on the nature of the products present in the solution to be treated. For example, steric hindrance and the electronic effects of the phosphorous ligands are factors influencing the removal of the metal.

The same Suzuki coupling was performed using different phosphorous ligands: three monodentate and three bidentate. For comparison purposes, scavenging screening was done by using the same two sets of conditions. No optimization was done to increase Silia*MetS* performance. By experience, using longer reaction times or higher temperatures will allow better results.



	SiliaMetS Scavenging Results with Monodentate Ligands (in %) with 4 equiv, 4 h						
	Triphenylphosphine [PPh <sub>3</sub> ]		Tri(o-tolyl)phosphine [P(otol) <sub>3</sub> ]		Tri-n-butylphosphine [PnBu <sub>3</sub> ]		
Silia <i>MetS</i>	00						
	22°C	60°C	22°C	60°C	22°C	60°C	
SiliaMetS Thiol	70	97	87	96	26	85	
SiliaMetS DMT	95	97	95	> 99	36	87	
Initial Pd Concentration:	27 ppm	n EtOAc	84 ppm in EtOAc 90 ppm in EtOAc				

Silia/MetS Scavenging Results with Bidentate Ligands (in %) with 4 equiv, 4 h							
	1,1'-bis(diphenylph [dp	osphino)ferrocene pp[]	ene 1,3-bis(diphenylphosphino)propane [dppp]		(+/-) BINAP		
Silia <i>MetS</i>							
	22°C	60°C	22°C	60°C	22°C	60°C	
SiliaMetS Thiol	50	69	75	90	31	56	
SiliaMetS DMT	14	22	95	98	41	64	
Initial Pd Concentration:	63 ppm i	n EtOAc	93 ppm i	in EtOAc	16 ppm	in EtOAc	

### Conclusion

In all cases, Silia*MetS* DMT and Thiol remained the best scavengers throughout the study, even though there is a variation in the nature of the ligand.

As expected, scavenging was more difficult with bidentate phosphine ligands. The best conditions were using 4 equiv for 4 h, at 60°C.



Removal of Pt

# Polyhedral oligomeric silsesquioxanes (*POSSs*) are promising building blocks for organic-inorganic hybrid materials

Polyhedral oligomeric silsesquioxanes (*POSSs*) are promising building blocks for organic-inorganic hybrid materials, yet they have rarely been studied because the synthesis of bifunctional POSS monomers is a very difficult one in general.

In this work, a novel molecular design of a POSS monomer has been proposed and it was shown that substituents on the completelycondensed or incompletely-condensed POSSs can be widely varied. Furthermore, the resulting polymers are suitable for optically transparent materials because of their high thermal stability and transparency. The properties were controlled by the substituents on the silicon atoms of the POSSs and comonomers. The platinum-catalyzed hydrosilylation polymerization of the bifunctional POSS monomers was performed with siloxane comonomers.

For the measurement of optical properties, the obtained polymers films were fabricated according to the solution-casting method (*refer to experimental protocol*), and were approximately 50 µm thick, measured with a thickness gauge. They showed high transparency in the 380 - 400 nm range.



**Experimental Protocol**: A polymer (0.1 g) was dissolved in  $CHCl_3$  (1 mL) in the presence of SiliaMetS Thiol, and the solution was filtered and cast onto a glass substrate.

Initial and/or residual metal content was not disclosed.

Naka, K. et al. J. Polymer Sci. Part A: Polymer Chem., 2018, 9, 4108-4112 Kyoto Institute of Technology, Sakyo-Ku, Japan





Removal of Ru

# Electrostatic self-assembly and covalent fixation (*ESA-CF*) process for the synthesis of linear and cyclic polymers and their topology effects on thin-film electron mobility

A pair of topologically contrastive, linear and cyclic polymers having densely appended perylene diimide (*PDI*) units have been prepared by means of an electrostatic self-assembly and covalent fixation (*ESA-CF*) process.

As a first step, a pair of linear and cyclic poly(sodium acrylate) of different segment lengths were prepared through an atom transfer radical polymerization, followed by the end-capping reaction and subsequent ring-closing metathesis in the presence of Hoveyda-Grubbs catalyst second generation.

Unreacted precursor as well as unassigned side products were removed by flash chromatography and Ru removal with Silia*MetS* DMT.





The electron-only device (*EOD*) measurement by using thin-film samples of a series of the Cyclic Poly(perylene Diimide acrylates), revealed consistently higher electron carrier mobilities in comparison with the corresponding linear counterparts, the linear Poly(perylene Diimide acrylates).

Initial and/or residual metal content was not disclosed.

Tenuka, Y. *et al. Macromolecules*, **2016**, *49*, 5831-5840 University of Calgary, Alberta, Canada Dalhousie University, Nova Scotia, Canada

#### Removal of Ru

### SiliaMetS vs other purification methods for ruthenium scavenging

The use of Silia*MetS* to remove ruthenium catalysts after a ring-closing metathesis (*RCM*) reaction is a very effective purification method. One of its main advantages is that no product is lost during the purification step.

		EtO <sub>2</sub> C CO <sub>2</sub> Et	Degassed DCM Grubbs 2 <sup>nd</sup> Gen.	EtO <sub>2</sub> C_CO <sub>2</sub> Et		
				quantitative yield		
	S	cavenging Results f	or Various Purificatio	on Methods* ( <i>in %</i> )		
Scavenger Filtration over packed bed of <sup>2</sup> Flash Purification					irification	
Scaveriging	SiliaMetS DMT <sup>1</sup>	Act. Carbon	Celite	Silica	Manual	SiliaSep Cart.
Ruthenium captation	93	73	24	58	70	73

<sup>1</sup> Using 4 equiv, 16 h, 22°C.

<sup>2</sup> Solution is passed directly on a packed bed of various adsorbents, which was then washed with the same quantity of solvent.

\* Quantitative yield obtained for each purification method (*adjusted in function of the residual concentration of catalyst*). No impurities were generated in all cases using the different methods (*determined by NMR*).





# Photoinduced topological transformation of cyclized polylactides for switching the properties of homocrystals and stereocomplexes

Cyclized poly(L-lactide) (*PLLA*) and poly(D-lactide) (*PDLA*) incorporating an o-nitrobenzyl group as a photocleavable linker were synthesized and photoirradiated for topological transformation to form photocleaved linear polylactides.

The o-nitrobenzyl group was chosen as it is one of the most widely used photolabile protecting groups due to its prompt removal.

Two cyclic PLLA and PDLA with this photocleavable linker were prepared via a Hoveyda-Grubbs catalyzed olefin metathesis, a polymerization known to form highly optically pure polylactides without degradation in the stereochemistry. Silia*MetS* DMT was used to remove residues of ruthenium.

It was shown that the enthalpy of the photocleaved cyclized polylactides was much lower than that of the linear prepolylactides, hence the crystalline-amorphous interface is more stable for the cyclized polylactides arising from the topology.



#### Removal of Ru

# Regioselective ring-opening metathesis polymerization of 3-substituted cyclooctenes

Allyl-substituted cyclooctenes with various sidechains [amides, methoxy, methoxy-terminated oligo(ethylene glycol)s, and tetrahydrofurfuryloxy groups] were prepared as monomers and polymerized by ring-opening metathesis polymerization (*ROMP*) using Grubbs second-generation catalyst.

In all cases, the ROMP of allyl-substituted monomers proceeded in a regio- and stereoselective manner to afford polymers with remarkably high head-to-tail regioregularity with high trans-stereoregularity.

Poly(3ROCOE) were purified from contaminating Ru with SiliaMetS DMT (O.N. at 40°C)





# An Idenix case study: ruthenium removal in the macrocyclization of dienylureas via RCM, using Silia*MetS* DMT

A novel assembly of two structurally related 14-membered ring macrocyclic hepatitis C virus protease inhibitors was reported by Idenix Pharmaceuticals. Key to their successful construction was an ultimate ring-closing metathesis step on the highly functionalized dienyl-urea via Zhan Catalyst-1B (*Ru-based catalyst*).



Several methods have been reported to remove Ru by-products, and were investigated in this study with some variations, including the use of tris(hydroxymethyl)phosphine, lead tetraacetate, TPPO, DMSO followed by silica gel filtration, adsorption onto silica gel, activated carbon and silica gel chromatography, treatment with mercaptonicotinic acid (*MNA*) and washing with aqueous NaHCO<sub>3</sub>, and the use of supercritical fluid extraction. Resulting Ru levels of those methods vs Silia*MetS* DMT treatment were analyzed by ICP-OES and are listed below.

	Effect of Reaction Conditions and Purification on Ru Content and Yield of Protease Inhibitor							
Entry	Reaction Conditions	Ru Reduction Operation	Ru content (ppm)	Yield (%)				
		MNA / NaHCO $_{\rm 3}$ wash ; charcoal ; silica gel filtration	14 (initial)	-				
1	250 mL/g diene 4.9 mM	1 <sup>st</sup> MeOH trituration	12	63				
		2 <sup>nd</sup> MeOH trituration	7.5	58				
		Charcoal ; silica gel filtration	120 (initial)	-				
2	250 mL/g diene 4.9 mM 1.5 wt % catalyst	1 <sup>st</sup> MeOH trituration	34	81				
		2 <sup>nd</sup> MeOH trituration	20	75				
		Charcoal ; silica gel filtration	120 (initial)	-				
3	250 mL/g diene 4.9 mM 1.5 wt % catalyst	1 <sup>st</sup> MeOH trituration	48	79				
		5/4 v/v DCE / MeOH crystallization	4.6	61				
		Charcoal ; silica gel filtration	880 (initial)	-				
4	80 mL/g diene 15.2 mM	1 <sup>st</sup> MeOH trituration	300	84				
		Toluene crystallization	22	45				
		Charcoal ; silica gel filtration	880 (initial)	-				
5	80 mL/g diene 15.2 mM 1.1 wt % catalyst	1 <sup>st</sup> MeOH trituration 2:1 v/v	300	84				
		EtOAc / n-heptane crystallization	19	51				
6	80 mL/g diene 15.2 mM		380 (initial)	-				
0	1.25 wt % catalyst		66	75				
7	_ 160 mL/a diene 7.6 mM	CilioMote DMT 16 h tractment + filtration	200 (initial)	-				
1	1.25 wt % catalyst		3.6	86				
0	250 mL/g diene 4.9 mM		180 (initial)	-				
8	1.25 wt % catalyst		7.7	89				

### Conclusion

As one can easily note, treatment with Silia*MetS* DMT gave a much lower Ru residual content conjointly with the highest final yields. Hence, low yield losses (*11 - 14 %*) and high purities (*98 %*) were achieved, together with excellent Ru levels observed (*as low as 3.6 ppm*), indicating the strong utility of this approach in Ru removal in the synthesis of macrocycle HCV PIs IDX316.

B. A. Mayes *et al.*, *Org. Proc. Res. Dev.*, **2013**, *17*, 811-828 Idenix Pharmaceuticals Inc., Massachusetts, United States





### **Ruthenium scavenging**

Ruthenium-based catalysts are commonly used in organic synthesis, mainly in olefin metathesis reactions [*ROM(P*) and *RCM*]. Grubbs and Hoveyda-Grubbs catalysts are the most popular ruthenium-based complexes in this field of application. Complete ruthenium removal can be tedious using conventional methods.

Under various conditions, several Silia*MetS* are known to be excellent scavengers to obtain minimal tolerated concentrations of residual ruthenium.

A ruthenium scavenging study was conducted and various parameters were investigated in order to learn more about their influence on the scavengers' robustness as well as to establish the best experimental conditions.

Ruthenium Scavenging Results using SiliaMetS (in %)											
SiliaMats	Grubbs 1 <sup>st</sup> Gen.		Grubbs 2 <sup>nd</sup> Gen.		Hoveyda-Grubbs 1 <sup>st</sup> Gen.		Hoveyda-Grubbs 2 <sup>nd</sup> Gen.				
Siliamets	Toluene <sup>1</sup>	DMF <sup>2</sup>	Toluene <sup>1</sup>	DMF <sup>2</sup>	Toluene <sup>1</sup>	DMF <sup>2</sup>	Toluene <sup>1</sup>	DMF <sup>2</sup>			
Silia <i>MetS</i> Thiol	90	96	-	99	97	93	-	-			
SiliaMetS Thiourea	-	98	-	96	97	98	-	-			
Silia <i>MetS</i> DMT	95	99	> 99	99	> 99	98	98	99			
SiliaMetS Diamine	99	99	91	94	> 99	98	-	90			

**Notes:** Silia*MetS* Cysteine and Imidazole were not screened in this study. Only Silia*MetS* results higher than 90 % are presented in this table.

### **Experimental Conditions**

Nb. Equivalent: <sup>1</sup> 8 equiv of SiliaMetS, 16 h, 80°C

<sup>2</sup> 4 equiv of Silia*MetS*, 16 h, 80°C

Initial Concentration: 500 ppm for all ruthenium-based catalysts

#### Conclusion

For all Ru-catalyzed reactions, the best scavenging was achieved using Silia*MetS* DMT. For most experiments, little or no difference was observed when using toluene or DMF.

Removal of Sn

### Tin scavenging using SiliaBond Carbonate vs SiliaMetS Cysteine

SiliCycle also provides a quaternary ammonium salt grafted on silica to which a carbonate group is ionically bounded. The latter has shown to be an excellent alternative for tin retrieval from organic mixtures.



### **Stille Reaction**

Tin scavenging was demonstrated on a Stille coupling in which  $Bu_3SnBr$  is the major tin by-product. In each test, residual Pd was scavenged in its entirety (from 24 mg/L to < 0.1 mg/L).

Ya.		
Scavenger	4 h, 8 equiv, 22°C	4 h, 16 equiv, 22°C
Silia <mark>Bond</mark> Carbonate	91	99
SiliaMetS Cysteine	77	99

### Conclusion

Scavenging yields were excellent with both scavengers. Hence, Silia*Bond* Carbonate was found to be of the same high efficiency as Silia*MetS* Cysteine to scavenge tin compounds of R<sub>3</sub>SnX type.





Removal of Sn

### Tin acetate-promoted Buchwald-Hartwig couplings of heteroaromatic amines

Regioselective manipulation of hydroxyl groups in polyols has been critical to carbohydrate chemistry.

A robust selective allylation of 1-methyl-L- $\alpha$ -rhamnose was developed using n-Bu<sub>2</sub>SnO as the catalyst and proton sponge as optimal base for high regioselective control. The optimized condition afforded the 3-O-allylated rhamnose in excellent regioselectivity (> 20:1) and 82 % isolated yield on a 50 g scale.

1) nBu<sub>2</sub>SnO (0.2 - 0.5 equiv) RBr (1.1 - 2.0 equiv) TBAB (0.5 equiv) Proton Sponge (1.2 equiv) CH<sub>3</sub>CN, 69°C

2) SiliaMetS Cysteine (20 % wt) OR SiliaMetS TAAcONa (20 % wt)



83.8 - 96.1 % conv 17:1 - 34:1 A scalable isolation/purification process was developed which afforded the desired product in 72 % yield with 1530 ppm of Sn.

The residual Sn level can be further reduced to < 20 ppm by treatment with Silia*MetS* Cysteine & TAAcONa.

Resid	lual Sn ( <i>ppm</i> )	Sn ( <i>ppm</i> )				
Experimental Step	Starting Level	End Level	Recovery (%)			
Extractive Work Up	-	1,530	72			
1st treatment 20 wt % with SiliaMetS Cysteine	1,530	273	92			
2nd treatment 20 wt % with SiliaMetS Cysteine	273	31.9	93			
1st treatment 20 wt % with SiliaMetS TAAcONa	1,530	267	91			
2nd treatment 20 wt % with SiliaMetS TAAcONa	267	19.5	95			

also showed outstanding potential in the selective benzylation of the same substrate.

catalytic system

The isomeric purity remained unchanged after the scavenger treatment.

Li. X. et al. Org. Process Res. Dev., 2017, 21, 1653-1658

Process Development & Process Chemistry, DOW AgroSciences, Michigan & Indiana, United States

Removal of Sn

### Tin scavenging using SiliaMetS Cysteine & TAAcONa

The removal of tin residues can often be an issue due to the high toxicity of this metal. Traditional removal methods for this impurity are treatment with an aqueous solution of KF,  $NH_4OH$  or NaOH, or with bases such as DBU. However, the efficiency of these methods can vary and may be inapplicable for some particular compounds.

Both SiliaMetS Cysteine & TAAcONa can be used to efficiently remove tin residues from organic mixtures, as demonstrated below.



<sup>1</sup> Pd residues were completely removed after only one treatment with SiliaMetS Cysteine.



Removal of U

### Uranium XAFS analysis of kidney from rats exposed to uranium

The kidney is the critical target of uranium exposure because uranium accumulates in the proximal tubules and causes tubular damage, but the chemical nature of uranium in kidney, such as its chemical status in the toxic target site, is poorly understood. In this study, micro-X-ray absorption fine-structure (mXAFS) analysis was used to examine renal thin sections of rats exposed to uranyl acetate.

In the preparation of uranium standards for XAFS analyses, uranium adsorbed on 6 different metal scavengers were prepared: SiliaMetS TAAcOH, SiliaMetS Cysteine, SiliaMetS Amine, SiliaMetS DMT, SiliaMetS Imidazole & SiliaMetS Thiol.









resin were above 15,000 µg/g.



SiliaMetS Imidazole

The concentrations of uranium bound to the functionalized

SH

SiliaMetS TAAcOH

SiliaMetS Cysteine SiliaMetS Amine SiliaMetS DMT

SiliaMetS Thiol

Uranium Concentrations							
Scavenger	Uranium concentration ( $\mu g/g$ )						
SiliaMetS TAAcOH	149,000						
SiliaMetS Cysteine	77,640						
SiliaMetS Amine	37,400						
SiliaMetS DMT	87,530						
SiliaMetS Imidazole	74,400						
SiliaMetS Thiol	22,500						

National Institute for Quantum and Radiological Science and Technology, Chiba, Japan Japan Synchrotron Radiation Research Institute, Hyogo, Japan

Homma-Takeda, S. et al., J. Synchrotron Rad., 2017, 24, 456-462





### Enantiomer N-1 substituted tetrazole prodrug synthesis

The authors of the study performed a regioselective tin-mediated alkylation to access the N-1 alkylated tetrazole isomer. A highly selective enzymatic hydrolysis provided the desired prodrug enantiomer. For the final fragment union, a Suzuki-Miyaura coupling was employed, leading to the need to eliminate both tin and residual palladium catalyst from the API in multikilogram quantities.

Purification was done adding 10 kg of Silia*MetS* Thiol to 18.9 kg of the crude product solubilized in 2-MeTHF. The suspension was heated to 60°C for 2 h, then cooled to 2°C to isolate the compound. A 83 % yield was acheived with 99.1 % purity, having less than < 2 ppm of Sn and 9 ppm of Pd.









# A Genentech case study: palladium and ruthenium removal in the synthesis of AKT inhibitor Ipatasertib using Silia*MetS* Thiol (*multi-kilogram scale*)

The first-generation process to manufacture Akt inhibitor Ipatasertib through a late-stage convergent coupling of two challenging chiral components on a multi-kilogram scale was reported by Array BioPharma.

A carbonylative esterification and subsequent Dieckmann cyclization was developed to forge a cyclopentane ring in the target. A second key chiral component, a β2-amino acid, was produced using an asymmetric aminomethylation (*Mannich*) reaction.

Upon scale-up, the deprotection of the Boc-API for the preparation of the Ipatasertib mono-HCI salt was easily completed in toluene in 12 N HCI.

The aqueous layer was then basified to  $pH \ge 12$  with aqueous NaOH in order to extract the Ipatasertib free-base with DCM. The DCM solution was subsequently treated with charcoal and Silia*MetS* Thiol to remove colored impurities and trace heavy metals resulting from previous synthetic steps.

These metals consisted in Palladium (*Pd/C catalyst in a Noyori Asymetric Transfer Hydrogenation of ketone*), Ruthenium [(R,R) MsDPEN-Ru(p-cymene)Cl) catalyst for asymmetric ketone reduction] and Titanium (*TiCl*<sub>2</sub> catalyst in asymmetric Aldol addition).



In Akt inhibitor synthesis, Genentech describes a scalable catalytic asymmetric hydrogenation process for the multi-kilogram scale production of a  $\beta$ 2-amino acid, the last building block. An extensive catalysis screening and optimization study was done and identified a simple Ru-BINAP catalyst system to directly afford the S product in high enantiomeric excess and yield was reported.

The final process enabled the multi-kilogram production in > 99 % ee to be used as a key component for one of their clinical candidates.



#### Process Mass Intensity & Kg Solvents Input / Kg of $\beta\text{2-amino}$ acid

#### Conclusion

For 138 kg of the crude amino acid, 8.3 kg of Silia*MetS* Thiol were necessary. Filtration of the DCM solution over Celite<sup>®</sup> resulted in a 99 % yield of Ipatasertib free-base with a ruthenium content of less than 5 ppm (*ICP-OES*).

J. Lane, T. Remarchuk *et al.*, *Org. Proc. Res. Dev.*, **2014**, *18*, 1641-1651 Small Molecule Process Chemistry, Genentech, Inc., a member of the Roche Group, California, United States Array BioPharma Inc., Colorado, United States

T. Remarchuk *et al.*, *Org. Proc. Res. Dev.*, **2014**, *18*, 1652-1666 Small Molecule Process Chemistry, Genentech, Inc., a member of the Roche Group, California, United States Array BioPharma Inc., Colorado, United States

T. Remarchuk *et al.*, *Org. Proc. Res. Dev.*, **2014**, 18, 135-141 Small Molecule Process Chemistry, Genentech Inc., A member of the Roche Group, California, United States Catalysis and Chiral Technologies, Johnson Matthey, Cambridge United Kingdom WuXi AppTec Co., Ltd., Shanghai, China





# A Johnson & Johnson case study: Sonogashira reaction & metal scavenging of various metals

Johnson & Johnson, in collaboration with Solvias, published a paper in which a mild Sonogashira reaction was developed using various metal catalysts. Treatment with Silia*MetS* Thiol simultaneously removed Pd, Cu & Al. Residual concentrations were below 50, 10 and 3 ppm respectively, in the isolated product 3.



Note: copper comes from a previous synthesis step.

I. N. Houpis *et al.*, *Org. Proc. Res. Dev.*, **2009**, *13*, 598-606 Johnson & Johnson PRD, API Development, Belgium, and Solvias A.G., Synthesis and Catalysis, Switzerland





# A Pfizer Global R&D case study: palladium and copper removal on a large scale batch using Silia*MetS* Thiol

Pfizer published a paper on removal of palladium & copper impurities from a 20 kg pilot-plant batch. They made two subsequent treatments using Silia*MetS* Thiol (20 % + 7 % w/w) at room temperature for 12 hours. After scavenging with Silia*MetS* Thiol, the desired product was obtained with a yield of 76 %, containing only 17 ppm Pd and 1 ppm Cu.



An alternative method was also tried using 80 % w/w of Deloxan THP (DegussaAG) overnight followed by basification with Na<sub>2</sub>CO<sub>3</sub>. Residual metal concentration with this method was higher compared to that of Silia*MetS* and the yield was lower.

Screening Validation Results on Top 4 Scavengers										
Scavengers	Residual Metal Concentration (ppm) Pd Cu									
Degussa Deloxan THP	20	2	60 - 70							
SiliCycle SIliaMetS Thiol	17	1	76							

Silia*MetS* allows lower residual metal concentration & higher yield with fewer manipulations!

R. L. Dorow *et al.*, *Org. Proc. Res. Dev.*, **2006**, *10*, 493-499 Pfizer Global Research and Development, Michigan, United States



### Multiple metal scavenging

SiliaMetS can be used to remove multiple metals in the same reaction with excellent efficiency.

PdCl<sub>2</sub> (dppf).DCN THF

The Negishi coupling presented in scheme above was performed to show that Silia*MetS* can be used to simultaneously remove residual palladium, iron and zinc present after the reaction.

Multiple Removal Scavenging Results (in %)											
SiliaMetS	Pd	Fe	Zn								
SiliaMetS Cysteine	95	> 99	98								
SiliaMetS DMT	83	93	99								
SiliaMetS Imidazole	84	91	97								
SiliaMetS TAAcONa	97	> 99	> 99								
Initial Concentration (in THF)	188 ppm	110 ppm	6 ppm								

Conditions: 4 equiv of SiliaMetS (relative to palladium), 4 h, 22°C.

For Zinc removal, all tested scavengers gave excellent results. Overall, for multiple removal scavenging results, Silia*MetS* Cysteine and TAAcONa showed to be the most versatile scavengers.





# Scavenging of Pd(dtbpf) in Suzuki coupling for the synthesis of a DGAT-1 inhibitor

Diacyl glycerolacyltransferase-1 (*DGAT-1*) is one of two known isoforms that catalyse the final step of triglyceride biosynthesis and hence could play a role in the development of obesity and insulin resistance.

In a DGAT-1 inhibitor synthesis project, a kilogram-scale Suzuki-Miyaura reaction was described by Abbott researchers in 2010.

This synthesis was reproduced in SiliCycle's labs and the resulting contaminated mixtures were treated with our most promising scavengers.



Ef	Effect of Time in the Scavenging Efficiency with 4 equiv. at 22°C (in %)								
Securemente / Conditione	1	h	4	h	16 h				
Scavengers / Conditions	Pd	Fe	Pd	Fe	Pd	Fe			
SiliaMetS Thiol	53	19	72	31	90	67			
SiliaBond Tosic Acid	11	54	24	92	22	95			

89%

Effect of Time in the Scavenging Efficiency with 8 equiv. at 22°C ( <i>in %</i> )									
Scavengers / Conditions	1	h	4	h	16 h				
	Pd	Fe	Pd	Fe	Pd	Fe			
Silia <i>MetS</i> Thiol	69	12	86	37	96	69			
SiliaBond Tosic Acid	24	91	33	91	17	92			

Effect of Time in the Scavenging Efficiency with 4 equiv. at 45°C ( <i>in</i> %) Treatment with 2 Scavengers Simultaneously: Silia <i>MetS</i> Thiol & Silia <i>Bond</i> Tosic Acid									
Scavengers / Conditions	1 h		4 h Pd Ee		16 h Pd Ee				
	Fu	FC	Fu	FC	Fu	FC			
SiliaMetS Thiol & SiliaBond Tosic Acid	78	81	91	81	96	81			

### Conclusion

In this case again, it was Silia*MetS* Thiol that proved to be the preferred scavenger for palladium under any conditions. As for iron, Silia*Bond* Tosic Acid was, again and by far, the most effective.

Scavenger combinations can achieve, in some cases, even higher removal of both metals. The best conditions were using 4 or 8 equiv at 22°C for 16h.

M. Ravn et al., Org. Proc. Res. Dev., 2010, 14, 417-424
 D. Barnes et al., Org. Proc. Res. Dev., 2009, 13, 225-229
 Abbott Laboratories, North Chicago, Illinois, United States





### Scavenging of Pd-118 using SiliaMetS metal scavengers

Some of Silia*MetS* Metal Scavengers have a particularly powerful scavenging behavior toward reagent Pd-118.

Pd-118, or  $Pd(dtbpf)Cl_2$  is a strong, homogeneous catalyst that has been shown to be very stable and active for all coupling reactions, especially aminations, reductive carbonylations and Suzuki couplings. Nevertheless, metals from this active catalyst can act as severe contaminants and be tricky to get rid of.



Pd-118 Complex

### Behavior of Silia*MetS* Metal Scavengers Toward this Specific Complex (*catalyst only in solvent*)

Scavenging Efficiency of Preliminary Screening (in %)										
Convondoro	DMF, 22°C		DMF, 60°C		Methyl-THF, 22°C		Methyl-THF, 60°C		DCM, 22°C	
Scavengers	Pd	Fe	Pd	Fe	Pd	Fe	Pd	Fe	Pd	Fe
Silia <i>MetS</i> DMT	76	45	97	51	85	63	97	47	96	83
SiliaMetS Thiol	51	28	72	34	50	50	88	64	36	48
SiliaMetS Cysteine	59	49	67	49	70	60	93	52	53	50
Silia <mark>Bond</mark> Tosic Acid	25	47	8	54	70	69	85	83	80	78

**Conditions**: Palladium complex solutions were made in DMF, methyl-THF and dichloromethane, and shaken with 4 equiv. of metal scavengers for 4 hours, at room temperature or 60°C.

### **Scavenging Conclusion**

Silia*MetS* DMT and Cysteine proved to be the preferred scavengers for Palladium (*Pd*) in this application.

As for Iron (Fe), SiliaBond Tosic Acid (SCX) was the best scavenger.

Furthermore, very good removal of both metals could be achieved through a combination of both scavengers.







# Scavenging of Pd(dtbpf)Cl<sub>2</sub> in Suzuki coupling for the preparation of arylpyrrolidines

Abbott Laboratories has used  $Pd(dtbpf)Cl_2$  for the preparation of anylpyrrolidines resulting in a final mixture contaminated with Pd and Fe.



SiliCycle has used this latter synthesis as a control reaction to study the effect of our Metal Scavengers on such contamination. The final crude mixtures were treated with the most promising metal scavengers and further optimization was carried out.

# Overview of Metal Residues Scavenging Efficiency (*in* %) in Various Experimental Conditions of a Suzuki Coupling for the Preparation of Arylpyrrolidines

	Effect of Time in the Scavenging Efficiency with 4 equiv at 22°C (in %)								
Scavengers / Conditions	1	h	4	h	16 h				
	Pd	Fe	Pd	Fe	Pd	Fe			
Silia <i>MetS</i> DMT	78	44	89	51	98	35			
SiliaBond Tosic Acid	34	65	34	70	36	86			

	Effect of Time in the Scavenging Efficiency with 8 equiv at 22°C (in %)								
Scavengers / Conditions	1	h	4	h	16 h				
	Pd	Fe	Pd	Fe	Pd	Fe			
Silia <i>MetS</i> DMT	90	53	94	56	96	47			
SiliaBond Tosic Acid	31	85	32	89	33	91			

Effect of Time in the Scavenging Efficiency with 4 equiv at 45°C (in %)						
Scavengers / Conditions	1 h		4 h		16 h	
	Pd	Fe	Pd	Fe	Pd	Fe
Silia <i>MetS</i> DMT	81	57	90	58	95	34
Silia <mark>Bond</mark> Tosic Acid	27	81	28	85	29	89

#### **Scavenging Conclusion**

Clearly, Silia*MetS* DMT proves to be the preferred scavenger for palladium. As for iron, Silia*Bond* Tosic Acid was by far the best reagent. A combination of scavengers can achieve, in some cases, stronger removal of both metals. The best results were obtained using 8 equiv of each scavenger at 22°C for 16 h.

M. Ravn *et al.*, *Org. Proc. Res. Dev.*, **2010**, *14*, 417-424
 D. Barnes *et al.*, *Org. Proc. Res. Dev.*, **2009**, *13*, 225-229
 Abbott Laboratories, North Chicago, Illinois, United States



# SiliaMetS in aqueous conditions

Along with growing importance of sustainable chemistry and catalysis, Silia*MetS* compatibility in aqueous conditions needed to be evaluated. As a preliminary exploration, palladium nitrate scavenging was tested at four different pH in various acidic medias. In purely aqueous conditions, Silia*MetS* TAAcOH was the most compatible and efficient scavenger. However, in acidic conditions, apart from Silia*Bond* Tosic Acid, all scavengers showed good to excellent removal capability. Scavenging activity can either be driven by H<sub>2</sub>O+ concentration (*pH*), or its counter-ion.

Scavenging ( <i>in %</i> ) of Pd(NO <sub>3</sub> ) <sub>2</sub> in Various Aqueous Conditions, 4 equiv at 22°C, 4 h						
Scavengers	H <sub>2</sub> SO <sub>4</sub> (1 M)	HNO <sub>3</sub> (1 M)	AcOH (1 M)	H <sub>2</sub> O		
SiliaMetS Diamine	43	23	> 99	5		
SiliaMetS DMT	> 99	> 99	> 99	15		
SiliaMetS Imidazole	> 99	> 99	> 99	36		
SiliaMetS TAAcOH	98	98	98	98		
SiliaMetS Thiol	> 99	> 99	> 99	77		
SiliaMetS Thiourea	> 99	> 99	> 99	35		
SiliaBond Tosic Acid	19	6	76	28		
SiliaMetS Triamine	55	49	96	10		

Results illustrate well that the counteranion (*and counter-cation*) plays a determinant role in the affinity of the resin toward palladium.

**Experimental Conditions**: an aqueous 250 ppm solution of Pd(NO<sub>3</sub>)<sub>3</sub> was prepared in a volumetric flask. 8 mL of this solution was introduced in 10 mL polypropylene tube charged with four molar equivalents of a metal scavenger. All scavengers were treated identically. The tubes were shaken on the SiliCycle MiniBlock® orbital shaker for four hours. All solutions were filtered on separate tubes, and the remaining palladium was measured.

Other complexes were tested using the same method described as above for scavenging of Pd(NO<sub>3</sub>)<sub>3</sub>.

Various Metallic Complexes Scavenging ( <i>in %</i> ) & Concentrations in Aqueous Conditions, 4 Equiv at 22°C, 4 h						
Scavenger	RuCl <sub>3</sub> [150 ppm]	K <sub>3</sub> PtCl <sub>6</sub> [250 ppm]	FeCl <sub>3</sub> [250 ppm]	RhCl <sub>3</sub> [250 ppm]	Pd(NO <sub>3</sub> ) <sub>2</sub> [250 ppm]	
SiliaMetS Diamine	11	71	25	94	5	
SiliaMetS DMT	0	97	6	68	15	
SiliaMetS Imidazole	0	91	6	59	36	
SiliaMetS TAAcOH	63	0	> 99	5	98	
SiliaMetS TAAcONa	47	87	98	7	77	
Silia <i>MetS</i> Thiol	0	57	7	0	35	
SiliaMetS Thiourea	0	92	9	34	28	
Silia <mark>Bond</mark> Tosic Acid	52	87	> 99	98	99	
SiliaMetS Triamine	14	61	13	92	10	

Various Metallic Complexes Scavenging ( <i>in %</i> ) & Concentrations in Aqueous Conditions, 4 Equiv at 22°C, 4 h						
Scavenger	AgNO <sub>3</sub> [250 ppm]	Ni(NO <sub>3</sub> ) <sub>2</sub> [250 ppm]	Sn(OTf) <sub>2</sub> [250 ppm]	CuSO <sub>4</sub> [250 ppm]	ZnSO <sub>4</sub> [250 ppm]	
SiliaMetS Diamine	74	43	47	93	58	
Silia <i>MetS</i> DMT	> 99	40	60	86	51	
SiliaMetS Imidazole	90	25	39	64	39	
SiliaMetS TAAcOH	> 99	84	93	99	86	
SiliaMetS TAAcONa	97	96	73	94	95	
SiliaMetS Thiol	96	6	6	0	0	
SiliaMetS Thiourea	79	1	17	14	3	
SiliaBond Tosic Acid	86	90	95	88	81	
SiliaMetS Triamine	76	33	58	44	58	

### Conclusion

Silia*MetS* TAAcONa has shown to have very good scavenging efficiency in aqueous media. Silia*MetS* TAAcOH and Silia*Bond* Tosic Acid were also efficient. Generally speaking, many metal scavengers have demonstrated excellent performance in aqueous environment, even when bearing hydrophobic moieties.



# Application Notes & Case Studies Silia*Bond* Organic Scavengers

# Silia*Bond* trapping agent in the evaluation of tea tree essential oils as skin sensitizers

Authentic Tea Tree Oils (*especially aged ones*) have been identified as potential skin sensitizers. Thiols were used a trapping agents (*DCYA*) for isolation, characterization and identification of several possible DCYA-adducts, which can be used to deduce the structure of the candidate reactive species.

Silia*Bond* Maleimide was used to quench excess of thiols. Both Silia*Bond* Maleimide and Tosyl Chloride were tested, but Silia*Bond* Maleimide was found to be more selective in their experimental procedures.



III. Khan, I. et al. Toxicol. in Vitro, 2018, 46, 237-245

University of Mississippi, Mississippi, United States

Further studies on detection and identification of potential ingredients of concern for skin sensitization were later conducted using Silia*Bond* Maleimide as a thio-scavenger in HTS-DCYA assays.

## Proteome-wide profiling of clinical PARP inhibitors to reveal compoundspecific secondary targets



Poly(ADP-ribose) polymerase (*PARP*) inhibitors (*PARPi*) are a promising class of targeted cancer drugs, but their individual target profiles beyond the PARP family, which could result in differential clinical use or toxicity, are unknown.

Using an unbiased, mass spectrometrybased chemical proteomics approach, a comparative proteome wide target map of four clinical PARPi was generated. Silia*MetS* DMT and Thiol were useful as organic scavengers for three out of the four PARPi: Niraparib (*scheme 1*), Rucaparib (*scheme 2*) and Veliparib (*scheme 3*).

 Rix, U. et al. Cell. Chem. Biol., 2016, 23, 1490-1503
 H. Lee Moffitt Cancer Center & Research Institute, Tampa, United States University of California, Los Angeles, United States
 University of South Florida, Tampa, United States



# Aliphatic amines in Antarctic CR2, CM2, and CM1/2 carbonaceous chondrites amines

Meteoritic water-soluble organic compounds provide a unique record of the processes that occurred during the formation of the solar system and the chemistry preceding the origins of life on Earth. Hence, molecular, isotopic, and enantiomeric compositions of amines in Antarctic carbonaceous chondrites were studied.

Silia*MetS* DMT & Silia*Bond* Amine were used to purify the organic compounds from carbonaceous chondrites.



# A Broad Institute case study: ionic scavenging of acids with SiliaBond carbonate & scavenging of amines with SiliaBond carboxylic acid

In the context of the synthesis of a library of highly complex macrocycles with a pyran core, Silia*Bond* Carbonate (Si- $CO_3$ ) and Silia*Bond* Carboxylic Acid (Si-WCX) were used in the purification process.

In the reaction below, Silia*Bond* Carbonate was used to remove excess of the acidic acylating reagent, and used again in the next step to remove excess benzoic acid. Then, Silia*Bond* Carboxylic Acid was added to remove any excess N,N-Diisopropylethylamine or potential o-acylation by-products.

Purities of various macrocycles were between 70-92 %, with an average purity of 87 %.



### **General Procedures**

Acylation: The crude amino alcohols, cyano-fluorobenzoic acid 4-o or 4-p (0.14 mmol; 1.0 equiv), SiliaBond Carbodiimide (*Si-DCC*) (0.19 mmol; 1.4 equiv) and DIEA (0.090 mmol; 0.7 equiv) were combined in 2 % dimethylformamide (*DMF / DCM 3.0 mL*), and stirred at room temperature overnight. In cases where acylation is slow, additional Si-DCC (0.19 mmol; 1.4 equiv) and a solution of HOBt (0.03 mmol; 0.2 equiv) in DMF / DCM (1.0 mL) and DIEA base (0.03 mmol; 0.2 equiv) were added. After acylation was deemed complete, reactions were scavenged with Si-CO<sub>3</sub> (0.18 mmol; 1.4 equiv) and Si-WCX (0.18 mmol; 1.4 equiv) for 30 min and then filtered and evaporated for 4 h.

**SNAr Macrocyclization**: All crude products from above were dissolved in DMF (4.0 mL) and heated at  $110^{\circ}$ C with Cs<sub>2</sub>CO<sub>3</sub> (*approximately 0.61 mmol; 4.5 equiv for 4 h*). Reaction mixtures were filtered through Celite, washed with DCM, and solvents evaporated. Crude products were dissolved in DCM and treated with Si-CO<sub>3</sub> (0.18 mmol; 1.4 equiv) and Si-WCX (0.18 mmol; 1.4 equiv) for 30 min, and then filtered through Celite and concentrated.

E. Comer et al., Proceedings of the National Academy of Sciences of the United States of America, 2011, 108, 6751-6756 Chemical Biology Platform, Broad Institute, Cambridge, MA 02142



# An Abbott case study: ionic scavenging of boronic acids with Silia*Bond* carbonate

An efficient Suzuki coupling protocol was developed and excess boronic acids were rapidly removed using solid-phase extraction and Silia*Bond* Carbonate to yield clean product.



### **General Procedure**

A Smith Process vial was charged with a stir bar, 4-bromobenzonitrile, p-tolylboronic acid and 2 mL of ethanol. A solution of 1 M  $K_2CO_3$  was added followed by the Pd catalyst. The reaction vessel was sealed and heated to 110°C for 10 minutes under microwave irradiation.

After cooling, the reaction mixture was transferred to a pre-packed column of Silia*Bond* Carbonate which had been conditioned with MeOH /  $CH_2Cl_2$  (1:1), and the eluent was collected via gravity filtration.

The column was then washed with MeOH /  $CH_2Cl_2$  (1:1) (3 × 3 mL).

The eluents were combined, concentrated and purified by flash chromatography to yield a purified compound.



# A Roche case study: nucleophilic scavenging of acyl chlorides with Silia*Bond* Amine

The guanine synthesis started with the simple and direct acylation of O-benzyl-2-aminopurine with polymeric BEMP. This convenient acylation approach using polymeric base was attempted in order to avoid the laborious Mitsunobu reaction introducing N9 substituents and tedious aqueous work-up steps. Silia*Bond* Amine was used to sequester the excess acyl chloride instead of using polystyrene-based trisamine resins.



### **General Procedure**

Silia*Bond* Amine (1.5 equiv) was added to the reaction mixture and stirred for 1 h at room temperature. The Silia*Bond* Amine scavenger with bound acyl chloride was then filtered off and rinsed with solvent (*e.g.: MeOH*) to yield an acyl chloride-free solution.

K. Kim et al., Tett. Lett., **2000**, **41**, 3573-3576 Roche Research Center, Hoffmann La-Roche, Inc., Nutley, NJ, USA



# Ionic scavenging of phenols and acids with SiliaBond carbonate

The efficiency of Silia*Bond* Carbonate as a scavenger of various coupling reagents (HX) - including pentafluorophenol, N-hydroxysuccinimide (*HOSu or NHS*), 4-nitrophenol, 1-hydroxybenzotriazole (*HOBt*) and 1-hydroxy-7-azabenzotriazole (*HOAt*) - was studied, as well as a comparison with two suppliers of polymer-supported carbonate.



Phenol Scavenging Results (in %)						
нх	SiliaBond Carbonate		Polymer 1		Polymer 2	
	5 min	60 min	5 min	60 min	5 min	60 min
Pentafluorophenol <sup>1</sup>	98	98	92	95	85	94
N-Hydroxysuccinimide	93	> 95	41	64	40	42
4-Nitrophenol	94	96	89	95	77	88
1-Hydroxybenzotriazole <sup>2</sup>	88	96	68	92	26	96
1-Hydroxy-7-azabenzotriazole <sup>2</sup>	97	97	72	96	30	92

Initial concentration: 5,000 ppm - 3 equiv of SiliaBond Carbonate. Analyzed by UV. 1 Analyzed by GC-MS, 2 in THF.



Benzoic Acid Scavenging Results ( <i>in %</i> )					
нх	Yield	Purity			
No Catalyst	35	95			
N-Hydroxysuccinimide1	67	98			
1-Hydroxybenzotriazole <sup>2</sup>	99	98			
1-Hydroxy-7-azabenzotriazole <sup>2</sup>	100	99			

**Note**: 1.0 equiv of amine, 1.5 equiv acid, 1.7 equiv catalyst (HX), 2.0 equiv Silia*Bond* Carbodiimide, 7.0 equiv Silia*Bond* Carbonate. Yield refers to the mass of isolated product. Purity was determined by GC-FID. 1 in DCM, 2 in THF.

### Conclusion

For each of the various experimental conditions / coupling reagents that were tested, Silia*Bond* Carbonate yielded better to much better scavenging results than its polymer-bound Carbonate equivalents.

Silia*Bond* Carbonate was also very useful in the scavenging of benzoic acid, in the following amide coupling, using Silia*Bond* DCC as a coupling reagent.



# Nucleophilic scavenging of boronic acids with SiliaBond Diol, SiliaBond Carbonate and SiliaMetS DEAM

Boronic acids and their derivatives are one of the most widespread intermediates and reagents in organic and medicinal synthesis. On-the-market drugs have even been adding boron atoms to enhance compatibility, selectivity and potency to their target molecules.

Up to very recently, boronic acids had always been reported as lacking apparent toxicity, mutagenic activity or in vivo instability issues. New studies have raised objections about this safe toxicological profile, both based on experimental and clinical data.

This is major in the chemical and pharmaceutical industry, as more and more studies denounce the genotoxicity of boronic acids. Today, in view of new data concerning boronic acids, there is pressure building on the ICH steering committee to assess boron compounds as potential genotoxic impurities (*PGIs*) per ICH M7.



All % of scavenging are determined by GC-MS

Depending on ones' needs, all three scavengers gave excellent scavenging results and showed to be an efficient and fast method for the removal of boronic acids.

Silia*MetS* DEAM and Silia*Bond* Diol are both excellent scavengers for Catch & Release: i.e., when the molecule of interest is temporarily bound either ionically or covalently to a functionnalized silica and subsequently released, once all undesirable impurities were washed out.



# Ionic scavenging of 2-iodobenzoic acid with SiliaBond TMA Acetate and SiliaBond Carbonate

Dess-Martin Periodinane (*DMP*) is a mild and chemoselective oxidant. It is readily accessible, environmentally benign and has a good shelf-life. Furthermore, the ease of handling, simple reaction work-up, product purification and good yields obtained with DMP make it a valuable reagent in organic synthesis.

2-lodobenzoic acid is the degradation product from DMP formed during the work-up. Most of it can be removed with a basic work-up, but sometimes, it can be difficult to get rid of all this side product.



### **General Procedure**

A solution of 1-octanol (1.00 mmol; 1.0 equiv) in  $CH_2CI_2$  (6 mL) at room temperature, was added to DMP (1.10 mmol; 1.1 equiv). The reaction mixture was stirred for 16 h, then diluted with 35 mL of MTBE and poured in 20 mL of an aqueous solution of Na<sub>2</sub>S<sub>2</sub>O<sub>3</sub> (25 %). The mixture was stirred for 10 min. Another portion of 35 mL of MTBE was added for the liquid-liquid extraction.

The MTBE phase was then washed with water and a saturated aqueous solution of NaCl (10 mL) and dried on MgSO<sub>4</sub>.

Scavenging was done using SiliaBond TMA Acetate or Carbonate, both in bulk (1 g) and SPE cartridge (6 mL / 1 g) for comparison purposes. Each sample was washed or eluted with a fresh portion of MTBE (8 mL) and then the 2-iodobenzoic acid concentration was monitored by GC-MS against an internal standard. The over-oxidation product (*carboxylic acid*) was succesfully scavenged using all products and formats.

Scavenging of 2-lodobenzoic Acid Results (in %)					
Scavenger	Bulk	SPE			
SiliaBond TMA Acetate	100	100			
SiliaBond Carbonate	100	100			

# Have a Scavenger Success Story?

Interested in sharing your scavenging application as one of our customer success stories? We would love to hear from you!

The get started, simply send us an email at <u>scavengers@silicycle.com</u> with your projects details. Details can include project title, challenges faced, project info such a procedure and results, and of course how Silia*MetS* made an impact. Bullet point is fine and the more details the better!

A member of our team will then contact you to get more information if needed and send over a first draft of your Case Study for you to review plus provide any feedback. Once you are satisfied and have approved the final draft, we will then share it for others to learn from!



# Carboxylic acids purification with SiliaBond TMA Acetate

Silia*Bond* TMA Acetate is an ion exchange media useful to extract organic anions from organic or inorganic matrices. It is less selective than Silia*Bond* TMA Chloride. The acetate anion being more labile than the chloride, it therefore retains more easily acidic compounds with pK<sub>a</sub> in the range of 4 - 5, such as carboxylic acids.

12 mL cartridges were filled with 2 or 4 g of SiliaBond TMA Acetate (*loading of 1 mmol/g*, for an equivalent of about 4 mmol of active function). They were tested with quantities of 1 and 2 mmol of each selected acid.

#### General Procedure for Catch & Release Purification

- 1. 12 mL cartridges (2 or 4 g of SiliaBond TMA Acetate) were conditionned with 6 mL of MeOH.
- 2. Compound was dissolved in 1 2 mL of MeOH and loaded onto the cartridge.
- 3. Column was washed with 15 mL of MeOH.
- Compound was released with a solution of AcOH / MeOH : 2/98.
- 5. Solvents were evaporated and final compound weighted.



#### 3-Chlorobenzoic Acid Purification Via Catch & Release



#### Ester Hydrolysis Purification Via Catch & Release



### Conclusion

Epoxidation reactions with 3-chloroperbenzoic acid (*mCPBA*) often yield after treatment of the reaction a certain undesirable amount of 3-chlorobenzoic acid, which can sometimes be difficult to separate from the desired product.

In the present example, 4-bromostyrene was treated with mCPBA, and the reaction medium is then treated with an aqueous solution of sodium sulfite. The latter allows to destroy excess reagent and eliminates much of the 3-chlorobenzoic acid correspondent. After extraction with dichloromethane and evaporation, the product was loaded on a 12 mL TMA Acetate SPE cartridge of 2 or 4 g. A simple elution with methanol made it possible to isolate 88 % of the desired epoxide product. Similarly, the same strategy was applied for the purification of 2-nitrobenzoic acid after hydrolysis of benzyl-2-nitrobenzoate, to yield the former molecule in a 90 % final yield.

In conclusion, Silia*Bond* TMA Acetate is very useful for the purification of carboxylic acids. Conversely, it may free the reaction media from compounds having a pKa lower than 5.





High Value Silica-based and Specialty Products for Chromatography, Analytical and Organic Chemistry, and Purification

### DISCOVER OUR BROCHURES



#### Overseas Offices

SiliCycle Europe SiliCycle India SiliCycle Shanghai

europe@silicycle.com india@silicycle.com contact@silicyclechina.com

SiliCycle 2022/06 EBK-SCAV001-0

< Follow us: in 🔰 F You Tube 🞯