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16 **Highlights**

- 17 A category of saturated alcohols was created
- 18 Data compilation was undertaken for the category of n-alkanaols
- 19 Repeat dose NOELs were read across for low toxicity compounds
- 20 *In vitro* data reduce uncertainty in read-across

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 Abstract: n-Alkanols provide an excellent example where a category-approach to read-across may be used to estimate the repeated-dose endpoint for a number of untested derivatives (target chemicals) using experimental data for tested derivatives (source chemicals). n-Alkanols are non-reactive and exhibit the unspecific, reversible simple anaesthesia or non-polar narcosis mode of toxic action in that the metabolic products of the parent alcohols do not contribute to the toxic endpoint evaluated. In this case study, the chemical category is limited to the readily bioavailable (C5 to C13) analogues. The toxicokinetic premise includes rapid absorption via the gastrointestinal tract, distribution in the circulatory system, and first-pass metabolism in the liver resulting in metabolism via oxidation to $CO₂$ and with minor elimination of oxidative intermediate as glucuronides. Two analogues have experimental 90-day oral repeated-dose toxicity data which exhibit qualitative and quantitative consistency. Typical findings include decreased body weight, slightly increased liver weight which, in some cases, is accompanied by clinical chemical and haematological changes but generally without concurrent histopathological effects at the Lowest Observed Effect Level (LOEL). Chemical similarity between the analogues is readily defined by a variety of structure-related properties; data uncertainty associated with toxicokinetic and toxicodynamic similarities is low. Uncertainty associated with mechanistic relevance and completeness of the read-across is reduced by the concordance of *in vivo* and *in vitro* results, as well as high throughput and *in silico* methods data. As shown in detail, the 90- day oral repeated-dose toxicity No Observed Effect Level (NOEL) value of 1000 mg/kg bw/d for 1-pentanol and 1-hexanol based on LOEL of very low systemic toxicity can be read across to fill the data gaps of the untested analogues in this category with acceptable uncertainty. **Keywords:** read-across, n-alkanols, repeated-dose toxicity, No Observed Effect Level (NOEL),

Lowest Observed Effect Level (LOEL), weight-of-evidence (WoE), uncertainty

1 Introduction

1.1 Read-across

 The principal philosophy of a toxicological read-across is chemicals that are similar in molecular structure will exhibit similar chemical properties, and as such, they will exhibit similar toxicokinetic and toxicodynamic properties. Thus, experimentally-derived toxicokinetic and toxicodynamic information and data from one chemical, the source substance, can be read across to fill the data gap for a second chemical, the target substance which is similar. This type of data gap filling is particularly useful for cosmetic ingredients where *in vivo* testing in Europe is prohibited by legislation [1]. As a predictive tool, read-across has been used by industry and regulators for decades [2]. With advances in non-animal test methods, read-across today is held to a different standard than at the turn of the century. Specifically, there is greater expectation in terms of the identifying similarities and addressing uncertainties within the read-across argument [3]. In order to facilitate the development of better practical guidance on how to formulate high quality read-across justifications, a series of case studies have been conducted by the authors. This case study illustrates specific considerations where metabolism of all the analogues in the chemical category is highly similar and plays no role in determining toxicological similarity [4]. The case study is also intended to illustrate how non-animal data, in the form of high throughput screening (HTS) data and *in silico* molecular screening, may be used to reduce uncertainties, as well as, add to mechanistic plausibility and weights-of-evidence (WoE) to any read-across argument.

 While it is easy to establish similarity based on structure and chemical properties, this similarity alone is often not enough to accept a toxicological read-across prediction for sub-chronic and

 chronic health endpoints. To justify the applicability domain of the category it is often necessary to establish toxicodynamic, and to a greater extent toxicokinetic, similarity within the category. The purpose of this research was to demonstrate the how read-across predictions of the repeated- dose toxicity no observed effect level (NOEL) value based on a consistent set of lowest observed effect level (LOEL) symptoms could be performed and substantiated for a category of n-alkanol analogues. Specifically, the category based data providing information to reduce uncertainties, and add to the WoE associated with read-across predictions of specified *in vivo* data. Thus, the estimations from the read-across are quantitative and with sufficiently low uncertainty that they may be used in risk assessments. As such, the predicted 90-day repeated-dose NOEL values are accompanied by sufficient relevant *in vivo* and non-animal test data to make the uncertainties equal to what would be expected from running a test using a protocol similar to Organization for Economic Co-Operation and Development (OECD) TG 408. In the present study, a previously reported 'strategy' was employed to assess similarities and overall completeness of the read-across [5].

1.2 C5 – C13 n-alkanols: overview of existing knowledge

 Historically, intermediate chain-length n-alkanols are considered nonpolar narcotics which act mechanistically in a manner similar to depressant anaesthetics. Fang, McKim, Koleva and their co-workers [6-8] reported multiple-regression type quantitative structure-toxicity relationships 87 (QSARs) for oral $log LDS0^{-1}$ data for rodents and the 1-octanol/water partition coefficient (log Kow). Comparison of measured toxicity data with predictions from baseline QSARs reveals that saturated monohydric alcohols consistently behave as classic nonpolar narcotics [9].

Due to bioavailability, and distribution and mechanistic considerations, the applicability domain

101 for this case study is limited to n-alkanols with a carbon atom (C) chain length range of C5 to

C13. For example, since longer-chain derivatives are typically transported via carrier molecules,

they are not included in this chemical category. Also, shorter-chain derivatives are not included

in this chemical category, as they have the potential to volatilise.

The general anaesthetic potency of several members of this homologous series of saturated

aliphatic alcohols was determined in tadpoles, using the loss of righting reflex as the criterion of

anaesthesia [13]. In this series, anaesthetic potency increased with chain length and was maximal

for 1-dodecanol. The cut-off in potency was between C12 and C14, such that 1-tridecanol was a

partial anaesthetic.

n-Alkanols within the range C5-C13 are expected to be readily absorbed by the gastrointestinal

tract and distributed in the blood in solution. n-Alkanols are metabolised mainly in the liver via

alcohol dehydrogenase to corresponding aldehydes and, subsequently, by aldehyde

dehydrogenase to the corresponding carboxylic acids [14]. The fatty acid derivatives of

intermediate size n-alkanols are readily taken up by mitochondria, where they are degraded by β-

- oxidation, especially in hepatocytes and myocytes [14]. However, generally <10% of the dose of
- these primary alcohols form glucuronic acid conjugates which are excreted in the urine [15].

 Voskoboinikova [16] and Opdyke [17] have summarised the historical literature on aliphatic alcohol toxicity. More recently, the toxicity of alkanols containing from one to six C-atoms has 119 been reviewed [18]. A cursory summary of the rat oral acute and oral repeated-dose toxicity of intermediate size n-alkanol are presented in Table 2. In general, n-alkanols acute oral toxicity 121 (i.e., LC50) is very low, ranging from 1500 to 5000 mg/kg bw with an average value of ≈ 3000 mg/kg bw. n-Alkanols are only slightly toxic in oral repeated-dose testing; typically, the rodent, oral, 90-day, repeated-dose NOEL in mg/kg bw/d is in the range of 1/2 - 1/3 the LC50 value. This value is characteristically based on clinical symptoms, haematological values outside the normal range, or whole body effects different from normal. However, if ingested in large enough quantities (i.e., near lethal doses), n-alkanols have the potential to cause systemic damage to the liver, heart, kidneys, and/or nervous system (see citations in Table 2 for details).

| Alcohol | Oral LD50 (mg/kg) | Reference | 90-d Oral NOAEL $(mg/kg$ bw/d) | Reference |
|----------------|----------------------|------------------|-----------------------------------|------------------|
| 1-Pentanol | 2200 | $[19]$ | 1000 | $[20]$ |
| | 3645 | $[21]$ | 1000 | $[21]$ |
| 1-Hexanol | 4590 | $[22]$ | 1127 M | $[23]$ |
| | 4870 | $[24]$ | 1243 F | $[23]$ |
| 1-Heptanol | 3250 | $[24]$ | >1000 | $[26]$ |
| | 6200 M | [25, 26] | | |
| | 5500 F | [25, 26] | | |

Table 2. Rat oral acute and repeated-dose toxicity of selected n-alkanols.

^a NOAEL value is recorded as experimental result, but the details in the report indicate that it is read across from 1-dodecanol (CAS 112-53-8). $(CAS 112-53-8)$. M- male, F- female

 $\frac{131}{132}$

133 **2. Method and Materials**

134 This evaluation of selected n-alkanols follows the workflow of Przybylak et al. [5]. It is in accord

135 with the guidelines proposed by OECD [34] and Schultz and co-workers [35]. *In vivo* data used

136 in the assessment were taken from the literature, including ECHA REACH Registered

137 Substances database [36]. Mechanistic relevance, as well as toxicokinetic and toxicodynamic

138 similarity of the category analogues was established using relevant non-animal data.

139 2.1 Target and Source Substances

140 In this case study, the analogues (listed in Table 3) include seven target and two source

141 chemicals; the latter, those with repeated-dose data derived from a 90-day OECD TG 408 assay,

142 are noted in bold print. This list is inclusive, as defined by the limitations of the applicability

143 domain. The analogues represents n-alkanols which are found in governmental or industrial

144 inventories (e.g., OECD High Production Volume Chemicals). Additional substance identifier

145 information, such as chemical structures and molecular formulas, are available in Table 1 of the

146 supplemental information.

147 **Table 3.** n-Alkanols considered as part of the chemical category for read-across. Compounds in 148 bold indicate the source substances.

2.2 Endpoint

 The NOEL for the 90-day rat oral repeated-dose is the single endpoint for which this category approach is applied. The 90-day oral repeated-dose data for 1-pentanol and 1-hexanol are particularly well-suited for read-across; the NOELs are based on experimental results from a 4- dose exposure scenario (0, <100, between 100 and 500, and ≥ 1000 mg/kg bw/d) following a standard test guideline (OECD TG 408) where the LOEL symptoms are reported. Moreover, there are supporting repeated-dose results for 1-heptanol, 1-undecanol and 1-dodecanol from OECD TG 422 studies, with the exposure durations for males being 28 days and for females 54 days.

2.3 Hypothesis of the category

The premise for this read-across case study is:

161 • n-Alkanols of intermediate chain length (i.e., C5 to C13) are direct acting toxicants (i.e.,

metabolic activation and detoxification is not a factor in toxicity) with a similar reversible

mode of action (i.e., non-polar narcosis or simple anaesthesia).

 for 13 weeks [20, 21]. The "no-outward-effect level" (assumed to be the NOEL) was 1000 mg/kg/day.

 In a non-standard rat oral repeated-dose assay similar to an OECD TG 408 assay, animals were exposed to 0.25% (based on nominal concentrations in the diet) and 0.50% for 13 weeks; 1.0% for 10 weeks, then 2.0% (week 11), 4.0% (week 12) and 6.0% 13 weeks of 1-hexanol [23]. The 190 NOAEL for 1-hexanol was determined to be \approx 1100 mg/kg bw/d (1127 mg/kg bw/d for male and

1243 mg/kg bw/d for female rats).

While the endpoint read across in this exercise is the 90-day oral repeated-dose NOEL, there is

also high quality repeated-dose toxicity NOEL/LOEL data for shorter duration studies (e.g.,

OECD TG 422). Since these data are both qualitatively and quantitatively similar to the 90-day

 data, they may be used as WoE and to confirm that all category members are within the endpoint domain.

1-Heptanol was administered orally to rats under OECD TG 422 and 0, 100, 300 and 1000

mg/kg bw/d [26]. No treatment related changes were noted for all parameters (e.g., biochemical,

haematological and clinical parameters, as well as body weight, food consumption and

neurobehavioral effects).

 Following OECD TG 422, oral repeated-dose toxicity of 1-undecanol in rats was evaluated at 202 doses of ≈ 0 , 100, 500, 2000 mg/kg bw/d [30]. A NOEL for systemic toxicity of 2000 mg/kg bw/d was determined in male rats, in the absence of toxicologically significant effects at any dose level.

3.4 Data matrices for assessing similarity

 In order for a read-across prediction to be accepted, there is the requirement to establish similarity between the source and target substance [5, 34, 35]. While structural similarity is a minimum, toxicokinetic similarity, especially for metabolism, and toxicodynamic similarity, especially in regard to mechanistic plausibility, is required for sub-chronic endpoints such as 90- day oral repeated dose-toxicity [5].

3.4.1 Structural similarity

 As demonstrated in Tables 1 and 3 of the supplemental information, all the n-alkanols included in the category are structurally highly similar. Specifically, they: 1) belong to a common chemical class, aliphatic alcohols and subclass, n-alkanols, and 2) possess common molecular scaffolding, a C-atom backbone with a straight-chain configuration. Structurally, the only variable is the length of the hydrocarbon backbone, C5-C13.

3.4.2 Chemical property similarity

 As demonstrated in Table 2 of the supplemental information, all the n-alkanols included in the category have many of their physico-chemical properties determined experimentally. Thus, when required calculated values, which are based on these measured values can be accepted with high confidence. Properties, with the exception of density and pKa, trend in value related to C-atom number within the scaffold. Specifically, all category members exhibit molecular weights from 88 to 200 g/mol. Hydrophobicity (as modelled by log Kow) increases with number of C-atoms from >1.0 to <6.0, vapour pressure and water solubility decrease with molecular size, melting 244 point and boiling point increase with molecular size, and density is constant at 0.8 ± 0.1 g/cm³. 245 Since there is no readily ionisable substituent the pKa is consistent at \approx 15.2.

3.4.3 Chemical constituent similarity

 As shown in Table 3 of the supplemental information, all the n-alkanols included in the category have common constituents in the form of: 1) a single key substituent, -OH, and 2) structural fragments, -CH³ and -CH2-.

3.4.4 Toxicokinetic similarity

 Limiting the range of C-atoms for the applicability domain reduced the impact of size on adsorption, distribution, metabolism and elimination (ADME). From a bioavailability standpoint, the analogues exhibit in *in silico* models linear trend with molecular weight. Such modelling reflects hydrophobic-dependent uptake.

 The toxicokinetic understanding of alkanols is reasonably complete despite the fact that the experimental data, as summarised in Table 4 of the supplemental information, are limited. Absorption, distribution and elimination are not considered factors in these predictions. For example, 1-octanol is rapidly absorbed after oral administration (i.e., bioavailability >80%). 1- 259 Octanol is excreted mainly as CO₂, and to a lesser extent as n-octyl glucuronide [17, 27, 37]. 260 Other n-alkanols exhibit similar toxicokinetics, with n-alcohols generally forming <10% of the dose as glucuronic acid conjugates and are excreted in the urine [15].

 It is generally accepted that, regardless of species, metabolism of n-alkanols is highly efficient and proceeds in a similar fashion [38]. Basically, there only degradative or detoxification pathways involved in the metabolism of n-alkanols. It is universality accepted that in the first step of the biotransformation, the alcohols undergo stepwise intracellular oxidation to the 266 corresponding carboxylic acids, followed by a stepwise C2 unit elimination via mitochondrial β-oxidation [38].

3.4.5 Metabolic similarity

 As demonstrated in Table 5 of the supplemental information, all of the category members undergo oxidation and hydroxylation in metabolic simulations. Briefly, mammalian catabolism of fatty acids, which most often takes place in the mitochondria, leads to the formation of acetyl- coenzyme A (CoA), enters the TCA cycle and reduces nicotinamide adenine dinucleotide (NADH) and flavin adenine nucleotide (FADH2) which are used by the electron transport chain 274 to produce ATP [14].

275 While other processes, including ω -oxidation and α -oxidation, are known to take place, β - oxidation is the most common catabolic process in n-alkanol metabolism. It is highly likely that 277 the n-alkanols included in the category will be nearly completely metabolised (i.e., >90%) via the tricarboxcylic acid (TCA) cycle. It is generally agreed that cytosolic fatty acids are activated for degradation by conjugation with CoA. β-Oxidation of saturated fatty acids consists of a recurring cycle of four reactions [14]. In acids with an even number of C-atoms, this cycling continues until two molecules of Acetyl-CoA are produced in the final reaction. Acetyl-CoA is available to be further metabolised in the TCA cycle. In acids with an odd number of C-atoms, the end product is propionyl-CoA, which must be converted to succinyl-CoA to enter the TCA cycle.

3.4.6 Toxicophore similarity

 The severe limitation of the structural domains sharply reduces the likelihood of differences in toxicophores between the target and source analogues. As demonstrated in Table 6 of the supplemental information, none of the n-alkanols included in the category are associated with any toxicophore based on *in silico* profilers within the OECD QSAR Toolbox V3.4.

3.4.7 Mechanistic plausibility similarity

 concentration, whereas 1-dodecanol caused a stimulation of the ATPase activity. All alkanols studied caused an increased fluidity of the membrane; however, changes in the membrane fluidity do not seem to be a pre-requisite of the ATPase inhibition [41].

 The Fish Acute Toxicity Syndrome (FATS) approach put forth by McKim et al. [7] has furthered our mechanistic understanding and the effects of intermediate chain saturated alcohols in fish more than anything else. The FATS approach is based on physiological response sets from spinally transected rainbow trout (*Oncorhynchus mykiss*) exposed to model chemicals. Briefly, *in vivo* biochemical and respiratory-cardiovascular responses were measured during lethal aqueous exposures; the responses and their interdependence formed a complex data matrix, with the best response variables for mechanisms of action being determined with multivariate statistics. The FATS for 1-octanol is characterised by a striking slow-down in all respiratory and cardiovascular functions [7] that makes it distinct from other modes of actions. The action of 1-octanol is consistent with depressant anaesthesia.

 The contributions of functional groups in acute rat oral toxicity have been calculated using alkanes as the baseline [40]. The toxic contribution of the OH group is -0.108. This situation (negative contribution to toxicity as compared to corresponding alkane) has not been observed in acute fish toxicity because the threshold of excess toxicity is too high to distinguish differences in toxicity. Critical body residues (CBRs) calculated from percentage of absorption and bioconcentration factors indicate that most of aliphatic alcohols share the same modes of toxic action between fish and rat. Specifically, fish and rat log (1/CBR) and number of alcohols are 1.65; 18 and 1.58; 348, respectively [40].

 In summary, there are several lines of evidence that support the contention that all the analogues within the domain act in a similar fashion and that fashion is not different from simple

anaesthesia or non-polar narcosis.

3.4.8 Other *in vivo* endpoint similarity

 In mammals, alkanols are considered baseline inhalation toxicants which model as simple narcotics [9]. Based on acute oral toxicity, n-alkanols belong to Category 4 which do not require a hazard label for acute oral toxicity. Their LD50 values are very low, typically ranging from 342 1000 to >5000 mg/kg bw with an average value of ≈ 3000 mg/kg bw (see Table 2). In mammals, mild to moderate sub-lethal toxicity from a single oral dose of intermediate size alkanols include general gastrointestinal symptoms (e.g., nausea, vomiting, abdominal cramps and diarrhoea) associated with irritation. High ingested doses (i.e., near acute lethal levels) can cause gastrointestinal haemorrhage and liver injury. For example, in the rat, the LD50 for 1-octanol is >5000 mg/kg [17]; the only symptoms of intoxication observed were moderately to severely ruffled fur and mild sedation. The symptoms had disappeared completely 24 hours later. The growth of the exposed animals was similar to that of the controls.

In fish, alkanols are considered to act via the nonpolar narcosis mode of action [42, 43]. Within

the USEPA DSSTox Fathead Minnow Acute Toxicity (EPAFHM) database, alkanols are

represented. They exhibit toxic potencies not statistically different from baseline predictions.

Because of concerns for aquatic toxicity, a large number of alcohols, especially saturated ones,

have been tested *in vitro* for cell population growth inhibition [44]. Structure-activity results

from *in vivo* and *in vitro* tests are highly consistent [45]. Briefly, from a structural standpoint, the

aquatic toxicity of alkanols is partition-dependent, regardless of endpoint being assessed.

Generally, for alkanol exposures in *in vitro* assays, results are attributed to unspecific

 interactions with biological membranes [11]; such effects are typically directly correlated with 1- octanol/water partition coefficients (c.f. [46]).

3.4.9 Relevant *in vitro* and *in silico* data

 In an effort to further support the mechanistic argument for this read-across information form two new methods were examined. Specifically, relevant HTS data in the form of ToxCast data [47, 48] and of *in silico* nuclear receptor binding predictions [49] were evaluated. Within the USEPA toxicity forecaster program (ToxCast) [50], data are available for the majority of the n- alkanol derivatives (see Table 8 of the supplemental information). Of the 711 possible assays that form the ToxCast scheme, 1-octanol, 1-undecanol, 1-dodecanol and 1-tridecanol have been evaluated in 602 of them. Additionally, 1-hexanol, 1-heptanol and 1-decanol have been assessed in about 250 assays. Lastly, 1-nonanol has been tested in 150 ToxCast assays. The number of active assays varies from none for 1-octanol to 25 for 1-undecanol and 30 for 1-tridecanol. Within ToxCast, the n-alkanols are among the "least promiscuous chemical classes"; < 2.74% of the ToxCast assays showing any activity up to highest concentration tested and none of the active assay are associated with specific bioactivity.

Only four non-specific cell viability qHTS assays within the Toxcast suite were positive for four

of the tested n-alkanol analogues; no assay exhibits activity for five or more of the category

analogues. Specifically, the Tox21_ELG1_LUC_Agonist_viability,

Tox21_TR_LUC_GH3_Antagonist_viability, Tox21_AhR_viability and

Tox21_Aromatase_Inhibition_viability show a positive response with four n-alkanols but there is

no consistency among which analogues are positive.

 Alkanols were screened with a variety of *in silico* profilers [49]. Specifically, profilers for nuclear receptor binding were run to identify potential binding to the following nuclear receptors; PPARs (peroxisome proliferator-activated receptors), AR (androgen receptor), AHR (aryl hydrocarbon receptor), ER (oestrogen receptor), GR (glucocorticoid receptor), PR (progesterone receptor), FXR (farnesoid X receptor), LXR (liver X receptor), PXR (pregnane X receptor), THR (thyroid hormone receptor), VDR (vitamin D receptor), as well as RAR/RXR (retinoic acid receptor/ retinoid X receptor). The evaluation of potential binding to the receptors is based on structural fragments and physico-chemical features that have been identified as essential to bind to these nuclear receptors and induce a response. As noted in Table 6 of the supplemental information, no potential receptor binding was predicted. It is worth noting that ToxCast also tested for all of these receptors, and all corresponding assays were negative. Taken collectively, the HTS and *in silico* findings are not inconsistent with the cited *in vivo* data. The premise that, oral repeated-dose toxicity of n-alkanols are considered to be nonpolar narcotics and act in a manner similar to depressant anesthetics is consistent with the ToxCast data and receptor binding simulations results which indicate no activity associated with a specific mode of action.

4. Statement of uncertainty

 The categorical assessments of uncertainties along with summary comments are presented in Tables 4 and 5. Briefly, chemical similarity is limited by chain length but has no impact on repeated-dose toxicity. Data uncertainty with the fundamental aspects of toxicokinetics is low. Regardless of the species of mammals, all such category members are judged to be readily absorbed orally and to have similar distributions; metabolised via oxidation to the acid

415 **Table 4.** Assessment of data uncertainty and strength-of-evidence associated with the fundamentals of chemical, transformation/toxicokinetic and toxicodynamic similarity. 416 fundamentals of chemical, transformation/toxicokinetic and toxicodynamic similarity**.**

^aUncertainty associated with underlying information/data used in the exercise (empirical, modelled; low, medium, high)

- 419 high)
 420 ^bCons ^b Consistency within the information/data used to support the similarity rational and prediction (low, medium, high)
- 421
- 422
423

Table 5. Assessment of uncertainty associated with mechanistic relevance and completeness of

425 a Uncertainty: low, medium, high

426 **5. Statement of the conclusions**

427 This is the second in a series of read-across case studies; this particular study examines a

428 category of similar compounds that do not require (or do not undergo) metabolism to exert a

- 429 potential adverse human health effect [51]. *In vivo* oral repeated-dose exposure to n-alkanols
- 430 gives rise to a set of nonspecific symptoms, including clinical symptoms, haematological values
- 431 outside the normal range, or whole body effects different from normal. Limiting the category to
- 432 C5 to C13 analogues assures that the impact of bioavailability on the toxicokinetic and

 toxicodynamic profiles is very limited. Primary alkanols are direct-acting toxicants with a reversible mode of toxic action described as nonpolar narcosis (i.e., unspecific interaction with biological membrane in a manner similar to simple anaesthetics). The main route of exposure for alkanols is oral via rapid gastrointestinal absorption. The majority of an oral dose of any n- alkanol is promptly degraded via simple cellular oxidation; the remainder is eliminated as the glucuronide conjugate.

 Repeated-dose toxicity test results exhibit qualitative consistency in results between and within species. While protocols vary, results of oral repeated-dose testing exhibit qualitative consistency between and within mammals. Typical findings are only mild changes including decreased body weight, slightly increased liver weight, as well as clinical chemical and haematological changes, but typically without concurrent histopathological effects.

 Within ToxCast, the n-alkanols are among the "least promiscuous chemical classes"; < 2.74% of the ToxCast assays showing any activity and none of the active assay being associated with specific bioactivity. Screening with *in silico* profilers reveals that n-alkanols have no predicted potential of nuclear receptor binding.

This is a category read-across (i.e., many-to-one several times). While several analogues have

been evaluated experimentally in oral repeated-dose testing schemes, the 90-day oral repeated-

dose toxicity data and the NOAELs of 1000 mg/kg bw/d for 1-pentanol and1-hexanol is the

conservative prediction. A no systemic toxic conclusion with a NOAEL of 1000 mg/kg bw/d can

be read across with confidence to untested n-alkanols in the C5 to C13 category listed in Table 3.

6. Acknowledgements

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Supplementary material

Read-Across of 90-day Rat Oral Repeated-Dose Toxicity: A Case Study for Selected n-Alkanols

Tables for Assessing Similarity of Analogues and Category Members for Read-Across

| ID | Name | CAS No | SMILES | 2D Structure | Molecular Formula | |
|-------------------------|--------------|----------------|-------------------------------|---------------------|--------------------------|--|
| $\mathbf{1}$ | 1-Pentanol | $71-41-0$ | OH H_3C CCCCCO | | C5H12O | |
| $\overline{2}$ | 1-Hexanol | $111 - 27 - 3$ | CCCCCCO | H_3C | C6H14O | |
| $\mathbf{3}$ | 1-Heptanol | $111 - 70 - 6$ | CCCCCCCO | OH H_3C | C7H16O | |
| $\overline{\mathbf{4}}$ | 1-Octanol | 111-87-5 | CCCCCCCCO | H_3C , OH | C8H18O | |
| 5 | 1-Nonanol | 143-08-8 | CCCCCCCCCO | H_3C | C9H20O | |
| 6 | 1-Decanol | $112 - 30 - 1$ | CCCCCCCCCCO | HvC | C10H22O | |
| $\overline{7}$ | 1-Undecanol | 112-42-5 | H_2C CCCCCCCCCCCO | | C11H24O | |
| 8 | 1-Dodecanol | $112 - 53 - 8$ | CCCCCCCCCCCCO | | C12H26O | |
| 9 | 1-Tridecanol | 112-70-9 | CCCCCCCCCCCCCO | | C13H28O | |

Table 1: Comparison of Substance Identification, Structure and Chemical Classifications

 $M =$ measured value

¹Values typically derived from EPISuite v4.1, ^a KOWWIN Program (v1.68), ^b MPBPWIN v1.43, ^c at 25 deg C; (mg/L) Kow (WSKOW v1.42); ² ACD/Lab Percepta Platform - PhysChem Module (from ChemSpider); ³ Predicted by PERCEPTA; predicted by ACD (Advanced Chemistry Development Inc., Toronto, Canada)

| ID | Name | Key Substituent(s) | Functional Group(s) | Extended $Fragment(s)$ | Chemical Class: | Chemical Sub- Class: |
|-------------------------|--------------|---------------------------|-------------------------------|----------------------------------|------------------------------|---------------------------------------|
| $\mathbf{1}$ | 1-Pentanol | $-OH$ | $-CH_3$, $-CH_2$ | | saturated aliphatic alcohols | straight-chain |
| $\overline{2}$ | 1-Hexanol | $-OH$ | $-CH_3$, $-CH_2$ - | | saturated aliphatic alcohols | straight-chain |
| $\mathbf{3}$ | 1-Heptanol | -OH | $-CH_3$, $-CH_2$ | | saturated aliphatic alcohols | straight-chain |
| $\overline{\mathbf{4}}$ | 1-Octanol | $-OH$ | $-CH_3$, $-CH_2$ - | | saturated aliphatic alcohols | straight-chain |
| 5 | 1-Nonanol | -OH | $-CH3$, $-CH2$ | | saturated aliphatic alcohols | straight-chain |
| 6 | 1-Decanol | $-OH$ | $-CH3$, $-CH2$ | | saturated aliphatic alcohols | straight-chain |
| $\overline{7}$ | 1-Undecanol | -OH | $-CH_3$, $-CH_2$ | | saturated aliphatic alcohols | straight-chain |
| 8 | 1-Dodecanol | -OH | $-CH_3$, $-CH_2$ | | saturated aliphatic alcohols | straight-chain |
| 9 | 1-Tridecanol | $-OH$ | $-CH_3$, $-CH_2$ | | saturated aliphatic alcohols | straight-chain |

Table 3: Comparison of Substituents, Functional Groups, and Extended Structural Fragments

Table 4: Comparison of Abiotic Transformation and Toxicokinetics

^aWilliams, R.T. 1959. The metabolism of some aliphatic aldehydes, ketones and acids. In: Detoxication mechanisms. The metabolism and detoxication of drugs, toxic substances and other organic compounds, 2nd Ed., London: Chapman & Hall, Ltd., chapter four, pp. 88-113; bOpdyke, D.L. 1973. Monographs on fragrance raw materials. Food Cosmet. Toxicol. 11: 95-115; Kwok, E.S.C. and Atkinson, R., 1994. Gasphase atmospheric chemistry of dibenzo-pdioxin and dibenzofuran. Environ.Sci.Technol. 28:528-533; ^dAtkinson, R. 1994. Gas-phase tropospheric chemistry of organic compounds. J. Phys. Chem. Ref. Data, Monograph 2:1-216. Kamil, I.A., Smith, J.N. and Williams, R.T. 1953. Studies in detoxication. 46. The metabolism of aliphatic alcohols. The glucuronic acid conjugation of acyclic aliphatic alcohols. Biochem. J. 53: 129-136.

() - The number of metabolites for specific transformation.

Table 6: Comparison of Toxicophores

1 OECD QSAR Toolbox 3.3.² COSMOS profilers available via COSMOS space: http://cosmosspace.cosmostox.eu

Table 8: Comparison of Toxicologically Relevant *In Vivo, In Vitro* **and** *Ex Vivo* **Data**

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