About the Futile Dream of an Entirely Riskless and Fully Effective Remedy: Thalidomide

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Abstract: Hardly any other artificially made medicine has ever had such a dangerous and unexpected potential as thalidomide. The medicine, which started its life as a harmless sleep-inducing substance, caused neurological and teratogenic damage, and, at the moment is a promising candidate for the fight against leprosy, cancer, and AIDS. The present study focuses on the early German history. It takes the bioactivity of that remedy as a chemical property and supports the claim that if a substance is biochemically active, adverse effects can hardly be avoided. The ethical role of an ontological underdetermination of chemicals is discussed, and an analysis of particular responsibilities is presented along the lines of modern principlism in biomedical ethics.

Keywords: Applied ethics, Contergan, thalidomide, chemical ontology, teratogenicity, chemical underdetermination.

He who is at peace will not waver or become anxious by the constant changes in life, by the 'panta rhei' that is particular to our times. A harmless medicine ensures this steadfastness even in difficult situations.

1. Introduction

The Contergan disaster was possible because of the belief by laypersons and experts in the old myth of a remedy without any adverse effects, and the betrayal of that belief. Although the sleep-inducing sedative (or hypnotic) remedy Contergan – containing thalidomide as an active component – was officially launched by the company Grünenthal in Germany only in October 1957, the presumably very first victim of its adverse effects was born at Christmas of the preceding year. Ironically, this victim was the daughter of an employee of Grünenthal, who gave the substance to his pregnant wife,
apparently off the record. Their baby girl was born without ears (Stephens & Brynner 2001, p. 19), but the causal relation between the administration of the substance and this situation was far from obvious at that time. Many more humans – estimations go up to about 10,000 – were born suffering from similar malformations and several other conditions. And many more than 10,000 – perhaps five times more – died before any physician could explain why.

All this occurred around 1960, and the teratogen thalidomide was the substance that caused the catastrophe. The world was different after that substance had entered the scene.

‘What is a chemical substance?’ is a central question in the philosophy of chemistry, and it is just as central in ethics of chemistry, as well. This question is not sufficiently answered by pointing at the compositional or structural formula. All chemicals are long-term epistemic objects, meaning that the amount or realm of properties assigned to these substances can never be closed. Hence, a new substance always carries with it risks for living beings (Schummer 2001). Furthermore, regarding thalidomide, we have to take into account that it is not just one substance, but two (see Fig. 1).

Figure 1 Structural formula of the (R)-(+) enantiomer of thalidomide. The hydrogen atom at the asymmetric carbon atom points up, and the phthalimide group down. In the (S) enantiomer it is inverse.

This paper is divided into a narrative section and an ethical discourse section. According to its main purpose of introducing a pertinent case to chemists and chemistry students, the narrative shall focus on the main issues necessary for an ethical evaluation. As is appropriate in this specific case, the emphasis of study lies on the early (German) history up to about 1960. More elaborate exposures of the historical material and international aspects with partly different focuses can be found in the references, particularly in Sjöström & Nilsson 1972, Knightley 1979, Kirk 1999, Stephens & Brynner 2001, Zichner et al. 2005, and Lenhard-Schramm 2016.

Although there exist some cases of remedies with adverse effects in the history of pharmacy as, for instance, the salts of heavy metals like lead and mercury (and the alloys of the latter) and, more recently, the case of Vioxx; the case discussed here is unique in at least the following respects: the pre-
tended and offensively advertised harmlessness, the unexpected emergence of adverse effects (the malformational impact on new-borns and the damage to the neuronal system), the worldwide influence on the vigilance against drugs and drug safety regulations, and its huge catalytic effect on the development of neurology and teratology. Hence, thalidomide certainly ranks among the most influential man-made chemical substances of all time.

Section 2 narrates the emergence of the substance including a critical synopsis of the central (but questionable) scientific paper of Grünenthal scientists, the substance distribution into society, the contact of the drug with human beings, its chemical (stereoisomeric) peculiarity, its temporary extinction and proscription, and, briefly, its rebirth as an active medicine.

2. Main historical aspects

2.1 How did the compound come into existence?

From a chemical point of view, it seems likely that thalidomide was synthesized quite early, perhaps well before Grünenthal sold it as a remedy. The preparation path is not very complicated: Starting from glutamine or glutamic acid, the product can be obtained in two or three steps, respectively (Müller et al. 1999). However, astonishingly little is known about the early history of thalidomide. The customary or official version, here in the words of the pharmacologist Herbert Keller, who was closely involved, from his witness account at the German Contergan process (the ‘Alsdorf Trial’, 1968-1970), is the following:

During works in the field of derivates of glutamic acid, Doctor Kunz and I came across a new compound which was identified as N-Phthalylglutamic acid imide. Besides other pharmacodynamic effects I observed that the substance showed a, however not too impressive, sleep-inducing effect. [Cited after Kirk 1999, p. 52, my translation]

In their paper from 1956, Kunz, Keller & Mückter claimed that with respect to the increasing consumption of hypnotic medicines there emerged the need for effective drugs without adverse reactions. The most promising candidates seemed to be unsaturated tertiary alcohols on the one hand and piperidine derivatives on the other:

In our own work devoted to the preparation of peptides, 2,6-dioxo-3-amino-piperidines were synthesized. Their structural peculiarity [konstitutionelle Eigentümlichkeit] gave reason to prepare a number of such compounds and test them with respect to their efficacy in animal experiments. [Kunz et al. 1956, p. 426, my translation]
Of several prepared substances, ‘K 17’ (the 17th preparation Kunz made for Grünenthal, as the official version has it) seemed to be pharmacologically the most interesting one. However, none of the other substances are discussed in that paper (and have not been discussed since). K 17 is described as a white solid with a melting point of 271 °C. The solubility is severely restricted: K 17 is insoluble, for example, in water, methanol, ethanol, acetone, diethyl ether, and benzene, and it is only slightly soluble in dioxane and pyridine. Due to this, the administration of the substance was performed in the form of suspensions in most cases. The pharmacological effect was described as the ‘reduction of the motility’ of the animals shortly after administration, an impact contrasting with the effects of, for example, barbiturates and Doriden, which lead to an initial excitation. The main method the authors applied was the unusual ‘Zitterkäfig-Methode’ (jiggle-cage method): Eight male mice, respectively, were held in a cage hanging on a spring. The cage was connected to a platinum wire, which plunged into a bath of sulphuric acid, whenever the mice moved. On contact with the acidic bath the wire completed an electric circuit (24 V) and, through electrolysis, caused the formation of hydrogen which was measured volumetrically. By this procedure, during normal activity, the mice octets produced almost 6 ml/min H₂. When given the sleep-inducing drugs, the hydrogen volume was reduced. Being relevant as calibration standard, the authors defined the status ‘sleep’ by 50 % activity of the base level rather than measuring the hydrogen with actually sleeping mice. Apparently, it is almost impossible to induce sleep in eight mice at the same time in the same cage. Moreover, the dose-dependencies (and, thereby the determination of the smallest sleep-inducing doses) are not presented or discussed at all. Hence, this method is questionable because of major methodical drawbacks regarding calibration:

Logical enough, perhaps, this [the neglect of calibrationable test methods] led to the evolution of a test for sleep-inducing capacities in which there was no need for actual sleep to be produced. [Knightley 1979, p. 16]

Nevertheless, the paper was accepted and published, and the Grünenthal team came up with comparative results for six substances (Tab. 1), among them Doriden, the market competitor since 1955 (Fig. 2). According to the company’s goals, the most important conclusions from these data were the comparable efficacy of sleep-induction and the minimal toxicity.

The Sunday Times Insight Team gave the following interpretation of Table 1:

Without doubt this table is the origin of the illusion that thalidomide was something capable of overturning many decades of pharmaceutical experience: an effective drug and quite without side effects. [Knightley 1979, p. 18]
Table 1. Pharmacological/toxicological comparison of sleep-inducing drugs as inferred from the ‘jiggle-cage-method’ with mice (after Kunz et al. 1956, p. 428).

<table>
<thead>
<tr>
<th>Hypnotic Drug</th>
<th>Smallest Hypnotic Dose (mg/kg)</th>
<th>Start of Effect (min)</th>
<th>Duration (min)</th>
<th>Smallest Narcotic Dose (mg/kg)</th>
<th>Initial Excitation Phase</th>
<th>Lethal Dose LD50 (mg/kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thalidomide</td>
<td>100</td>
<td>5</td>
<td>240-300</td>
<td>&gt;5000</td>
<td>-</td>
<td>&gt;5000</td>
</tr>
<tr>
<td>Luminal</td>
<td>40</td>
<td>30</td>
<td>300</td>
<td>150</td>
<td>++++</td>
<td>300</td>
</tr>
<tr>
<td>Valamin</td>
<td>50</td>
<td>10</td>
<td>60</td>
<td>375</td>
<td>++</td>
<td>725</td>
</tr>
<tr>
<td>Doriden**</td>
<td>75</td>
<td>30</td>
<td>200</td>
<td>400-500</td>
<td>++++</td>
<td>600</td>
</tr>
<tr>
<td>Methyl-pentynol</td>
<td>150</td>
<td>15</td>
<td>210</td>
<td>500</td>
<td>++</td>
<td>750</td>
</tr>
<tr>
<td>Sodium Bromide</td>
<td>500</td>
<td>80</td>
<td>&gt;360</td>
<td>ca. 7000</td>
<td>-</td>
<td>8000</td>
</tr>
</tbody>
</table>

* The actual value of the p.o.-LD50 for rats is 113 mg/kg.
** Note that the competitor Doriden from Ciba is the only other substance of the ‘piperidine-family’ used here (see Fig. 2).

In the spring of 1955, clinical trials – the official testing of thalidomide on humans – began. No formal application was necessary at that time, and, moreover, no standard procedure for pharmacological trials had been developed yet. Although some side-effects were hinted at, all those reports concluded with similar positive judgments. Peripheral neuritis, in particular, was not described in detail as no thorough searches for side-effects had been conducted at the time, while the clinical studies used were far too short such that any long-term effects had insufficient time to develop. Thus, the severe adverse effects were found by epidemiological observations rather than in the clinical studies.

Figure 2. Molecular formulas of Doriden, thalidomide, and barbituric acid (from the left). Note the similar structure of the CO-NH-CO-sections of these molecules (also referred to as imide) which is considered one main locus for interactions with biological targets. Note also that thalidomide has two of these sections.
2.1 How did thalidomide get into the public?

With the backing by a study on nursing (but not pregnant) women (Blasiu 1958), thalidomide was advertised excessively. Two particular aspects were emphasized in this marketing campaign: The lack of toxicity and the natural-ness of the support of a ‘good sleep’. Furthermore, there seemed to be no risk of addiction. Generally, this led to a very comfortable description, which competitors like Doriden could by no means make a claim on (Kessel 2013). Physicians and apothecaries were the main marketing multiplicators and drug distributors for non-prescription drugs. Referring to the non-barbitural competitor Doriden of Ciba (see fig. 2), Nils Kessel describes the role of these experts as follows:

The key figure in this arrangement was the physician. He was analyzed as a potential consumer and a ‘consumption guide’ but officially treated as a partner in a common scientific endeavor. [Kessel 2013, p. 163]

In the case of thalidomide, recommendations (‘therapeutic circulars’) were distributed to about 50,000 experts, advising them to apply the drug in cases of sleep disorder, fear, contact weakness, nervousness, menopausal problems, female sexual excitability, and more. Obviously, marketing concentrated on female consumers. According to effective law, it was not allowed to address the lay public with advertisements on hypnotic drugs (Kirk 1999, p. 58). Nevertheless, the marketing campaign was very successful, and the revenues were remarkable. In the winter of 1960 Grünenthal made more than 1 million Deutsche Mark monthly with Contergan, an amount that decreased significantly one year later, when the drug was taken off the market. In 1960, however, those numbers represented almost 50% of the total turnover of the company. Clearly, the number of employees rose in parallel with the Contergan sales: from 420 in 1954, to 900 in 1959, and to 1300 in 1961. Understandably – from a plainly economical point of view – the company officials did not want to strengthen the impact of any possible critical statements in their ‘consumption guides’ or to introduce a prescription for their ‘wonder drug’. Yet in 1960, in an internal document from April 14, Mückter wrote:

Unfortunately, we are now receiving an increasing number of reports on the side-effects of this drug, as well as letters from doctors and pharmacists who want to put Contergan on prescription. From our side, everything must be done to avoid prescription enforcement, since already a substantial amount of our turnover comes from over-the-counter sales. [Cited after Sjöström & Nilsson 1972, p. 54]

Eventually, one year later, the management decided to apply for prescription requirement. However, the catastrophe could no longer be stopped.
2.3 How did it get banned?

Only when the previously mentioned presumed first victim of the malformational effects of thalidomide reached her fifth birthday did it become apparent that thalidomide was the cause for her having been born without ears, and for all the other victims. The main figures of this manifestation process were the physicians Widukind Lenz (1919-1995) from Germany and William Griffith McBride (b. 1927) from Australia. Apart from his professional work as a pediatrician and geneticist, Lenz (and his group) performed meticulous epidemiological studies and took the initiative to inform the learned public. He finally called Mücken by phone on 15 November 1961, but the latter was still reluctant to even admit his knowledge about reports on adverse effects. Today we know that Mücken was plainly lying. However, that would not help himself or Grünenthal in any way. On 26 November 1961, the German newspaper *Welt am Sonntag* published a long, scathing article about Lenz’s recent complaints entitled ‘Malformations from Tablets? Alarming Suspicion of a Physician against a widespread Drug’, and the next day Contergan vanished from the German market. In 1962, however, more than 900 new thalidomide victims were registered, and another 14 between 1963 and 1967. The following other countries reported thalidomide cases: Argentina, Australia, Austria, Belgium, Brazil, Canada, Denmark, France, Ghana, Great Britain, Israel, Italy, Japan, Lebanon, Luxembourg, Netherlands, Norway, Pakistan, Sweden, Switzerland, Spain, Syria, and USA. McBride’s letter to the editor of the scientific journal *The Lancet* (16 December 1961) is considered the very first publication on the embryopathic effects of thalidomide in humans. In that letter, McBride states that “congenital abnormalities are present in approximately 1.5% of babies,” and claims: “In recent months I have observed that the incidence of multiple severe abnormalities in babies delivered of women who were given the drug thalidomide (‘Distaval’) during pregnancy, as an antiemetic or as a sedative, to be almost 20%.” (McBride 1961)

The course of the (temporary) decease of thalidomide as a remedy was different in other countries. Even before Grünenthal’s withdrawal of the drug, the Richardson-Merrell Company failed to get approval to register thalidomide (under the brand name ‘Kevadon’) for commercial use in the United States, whereas in Canada the registration was successful. Frances Kathleen Oldham Kelsey (1914-2015), a physician and pharmacologist with the U.S. Food and Drug Administration (FDA), found the submitted material imperfect and unsatisfactory. Her main criticism referred to the obvious differences of the animal test results on the one hand and the impact on humans on the other. Kelsey remained unshaken in spite of heavy contentions with the officials of the company, postponed the approval repeatedly, and the company finally gave up the proposal in March 1962.
Intriguingly, the regulations in the United States and other countries like Germany allowed for ‘blind’ clinical studies on uninformed patients. That weak point in drug legislation lead to the circumvention of the Nuremberg medical code with its central claim of ‘informed consent’, which was compiled after the trials against the ‘Nazi doctors’ (Wiesing 2012, pp. 137-139). The obstetrician Ray Nulsen from Cincinnati, Ohio, for example, one of the large number of physicians Richardson-Merrell mentioned in their FDA application, repeatedly tested drugs for companies in this way. He published his experiences with thalidomide on pregnant women referring to ‘insomnia’ particularly during the third trimester which is just the period in which the fetus is insensitive to malformation (Nulsen 1961). This article was in fact written by Raymond Pogge, the medical director of Richardson-Merrell, rather than by Nulsen, and some of the important aspects were, to say the least, distorted. Even worse is that despite the published material and all his later testimony, Nulsen must have given the drug to women in earlier phases of pregnancy, too: More than a dozen cases of typical malformations and stillbirths he himself delivered are documented, and some of the former went to court. Most of these cases ended in the payment of compensation by Richardson-Merrell. We find a striking comment on the unfortunate role of Nulsen in that history, again by the Sunday Times Insight Team:

The final irony is that if Nulsen had been conducting a genuine clinical trial, or if Richardson-Merrell had been really interested in establishing the safe and effective application of thalidomide in a broad spectrum of circumstances, then they both might have won an honored niche in pharmaceutical history for being the first to discover thalidomide’s devastating effects on the unborn child. [Knightley 1979, p. 85, emphasis in original]

Hence, although thalidomide was not approved as a remedy in the USA, and therefore a disaster like in other countries did not emerge, even this country has experienced its pertinent cases as well.

Whereas their chapter about thalidomide in the United States carries the heading ‘The results of vigilance’, the Swedish authors Sjöström and Nilsson (both of whom have been active in the trials mentioned below) call the chapter about their own country ‘The results of negligence’ (Sjöström & Nilsson 1972). In Sweden, as opposed to (Western) Germany, Neurosedyn and Noxodyne, both containing thalidomide, were prescriptive drugs right from the beginning. Like Grünenthal, the company Astra too was very reluctant to withdraw its products. Because the reports about the withdrawal by Grünenthal referred only to the German trade name Contergan, both physicians and potential consumers in Sweden were perplexed and did not recognize possible risks. The result was that more than 100 Swedish children suffered from phocomelia (in a small country with about 7.5 million inhabitants). In February 1962, a few concerned parents founded an association which finally went
to court. Many international experts were invited. The issue ended in a settlement between the association for parents of malformed children and Astra. The compensation in Sweden (plus a few Danish cases) amounted to about 15 million USD.

2.4 The two faces of thalidomide

The purity of substances is a central and general topic in the history and philosophy of chemistry, but it becomes particularly intriguing in the instance of thalidomide, as it does for all stereo isomeric substances. Although all its molecules share the same compositional formula (and the same molecular spectra), the bulk material usually consists of two kinds of molecules which can be imagined like our two hands (see Fig. 1). The mixture of these enantiomers is called a racemate, and it is the latter that presumably was used in most experiments, particularly in those of the Grünenthal team, and presumably in the remedy, too.

The biochemical reactivity, however, depends on the particular spatial organization of molecules; in most stereoisomeric pairs we find different effects for each enantiomer (Knabe 1995). Taking into account their general indifference about the real properties of their 'wonder drug', there is no wonder that the Grünenthal researchers did not say anything about this issue. However, after the manifestation of the negative effects of thalidomide several research teams began investigations in that direction, such that thalidomide became the classical drug containing a 'bad' and a 'good' part. I briefly refer to that issue here. In 1965, R- and S- isomers were prepared (Shealy et al. 1965), and two years later, biochemists from London published the surprising result that all three entities, R-, S-, and racemic thalidomide, show a teratogenic impact on the New Zealand White Rabbit. This species had recently been specified to be the only species to react equally sensitively to thalidomide as humans do (and it is still used as an 'animal model' in teratogenicity tests during the pre-clinical development of new active substances). Even more surprising than the teratogenicity of any thalidomide species in rabbits was that a very persistent myth emerged and flourished after Köhler et al. (1971) had published their study about teratogenic effects on mice. According to this myth, the R(+) enantiomer is considered the sleep-inducing but otherwise harmless ('the good') one, and the S(-) enantiomer the embryotoxic ('the bad') one. The results of this study, however, which must rely on at least one misinterpretation, have been falsified by Scott et al. (1977), who clearly confirmed the missing teratogenic impact of thalidomide in rodent species other than rabbits. One important pharmacokinetical principle was – and still is – omitted or neglected by the followers of the myth of good/bad thalidomide: the metabolization of active compounds. In aqueous
systems like body liquids, thalidomide, although it is almost insoluble, can run through the well-known keto-enol tautomery, which is the dynamic joint between two different isomers. Only the keto-structure carries the asymmetry (Fig. 3).

The two enantiomers are transformed into one another, meaning they racemize. The half-life of both R- and S- forms in a neutral phosphate buffer is about five hours. The racemization is accelerated if enzymatic (catalytic) effects like serum albumin are present. With albumin, half-times are reduced to 18.5 min (R), and 9.5 min (S), respectively (Knoche & Blaschke 1994). Thus, both enantiomers are transformed/metabolized into the respective other form in the body, resulting after some hours in a racemic mixture. The differentiation of activities, as the myth has it, in fact cannot be tested empirically at all because of the peculiar nature of thalidomide. In addition to racemization, thalidomide is cleaved by hydrolytic reactions, such that it is still unknown which the active molecular agent is.

2.5 The unfinished story

Although it had a complicated and unclear beginning, was introduced to the public as a ‘wonder drug’, became a ‘dark remedy’, and has since been withdrawn as an active component, the epistemic and pharmaceutical object thalidomide is still vital and fashionable. At the moment, almost all of the
manifest effects including the inhibition of angiogenesis or endocrine disruption are far from being understood, but – or better because of this – the research is booming. The direction of research, of course, is turning away from the sleeping pill to the fight against leprosy, AIDS, and cancer. One of the most astonishing comebacks of a ‘dead remedy’, the healing of leprosy by thalidomide, has been initialized – by chance: In 1964, Jacob Sheskin (1914-1999), the director of the Jerusalem hospital, gave the medicine, which had already been withdrawn, to a hopelessly ill patient in 1964. After the intended good sleep, the pain and the leprosy symptoms almost immediately vanished. Many other patients reacted similarly. This unexpected positive impact was obtained by a desperate (‘blind clinical’) move. But still, although there might be some desirable effects with a drug like thalidomide, there will always be a remaining danger, namely the unconscious digestion of the substance. In countries like Brazil, where leprosy threatens a large number of people, unintended increases in the use of the substance has led to new cases of malformations. Therefore, as long as the substance is available to the public or parts of it, for whatever reason, vigilance must not be neglected (but this is already a point that belongs in the following section).

3. Ethical Discussion

Commenting on the behavior of the scientific decision-makers of companies like Grünenthal (Germany), Richardson-Merrell (USA), Distillers (UK), Astra (Sweden), Dainippon (Japan), etc., Roald Hoffmann writes:

[…] yes, this is abysmal science. And whereas science as a system for gaining reliable knowledge works in spite of instances of poor-quality experimentation – it will easily survive sloppiness, hype, and even fraud – the kind of science that touches on human lives cannot afford to be bad. The thalidomide disaster should not have been allowed to happen. [Hoffmann 1995, p. 136]

In fact, science survived these greed-driven distortions, lies, and slackness, and made significant progress, but many thousands of humans could not do so, and some thousands have suffered unnecessarily. Apparently, the transition from miserable science to ‘fraud and deceit in science’ is much easier than we would wish, and this has a reasonable influence on the ethical judgment, as we will see below.

3.1 Could the disaster have been avoided?

To start the discussion of pertinent ethical issues, that is to find out the neuralgic points of responsibilities, I shall explore some possible answers to the
question used as the heading of this section. A first, admittedly quite simplis-
tic answer would be: yes, if nobody had synthesized the substance, and if
nobody had distributed it, and if nobody had needed and consumed these
sleeping-pills during the first trimester of pregnancy, the surprising and
threatening chemical behavior would not have been realized – at least not in
the terrible way it was. A safety-oriented and less trivial positive answer could
be motivated by the evidence of neurological impairments, the reports of
which commenced in 1959. This less trivial ‘yes’ could be applied if physi-
cians and scientists, particularly those of Grünenthal, had been much more
attentive and careful than they were. The story of the non-licensing of tha-
ridomide to Richardson-Merrell in the USA due to the resistance of the FDA
physician Frances Kelsey (who in 1962 was honored by President Kennedy)
refers to that version to argue for a ‘yes’. Of course, Kelsey could not and
did not know that thalidomide would turn out to be a teratogen, thus that
part of the rejection of the application by the FDA was a lucky outcome.
Still, the officials of Grünenthal did a less than mediocre job as scientists and
physicians by dropping their scientific curiosity and veracity with respect to
their own product. While neglecting the underdetermined chemical character
of their product, they decided to follow or tolerate an aggressive marketing
campaign based on at least incomplete and distorted results and trivialized or
neglected all the anxious reports about side effects.

Another simple but naively whiggish justification of a ‘yes’ to the ques-
tion in the heading of this section would be connected to the performance of
teratogenicity tests. The problem is: nobody asked for such tests at that time
– not the governmental legislators or regulators, and in particular not the
manufacturers – and, furthermore, nobody would have known which experi-
ments to perform or which kind of animal to take. As described above, Blasiu
and particularly Nulsen came the closest to the pertinent knowledge, howev-
er by dubious methodologies. The missing-question-argument is supported
by the switch-on-effect of the number of publications after 1961 (for a few
German journals, see Fig. 4): Even the otherwise attentive scientific commu-
nity did not suspect anything like this, and a worldwide investigation cam-
paign with hundreds of papers was started only after the manifestation of the
effect by people like Lenz.

Finally, then, the answer to the question above must be ‘no’, if someone
would refer to the mentioned myth of the ‘evil’ and the ‘good’ thalidomide
(‘distomer’ vs. ‘eutomer’): there is – unfortunately, perhaps – no practical
way to differentiate those two biochemically.
3.2 Underdetermination – chemically

The chemical world is in flux. Every new substance can potentially come across many others – those already existing on the one hand, and those which come into the world later on the other. Hence it is impossible to give an exhaustive description of the chemical character and future behavior of chemical entities – in particular not by just pointing at simplifying compositional formulas (like some analytical philosophers still would like to have it). Even a long-standing and intensively researched substance such as water is not fully known up to the present. Chemical substances, even those that are assigned the status of a ‘natural kind’ by some authors (which is not applicable in the present context, because thalidomide clearly is an artificial product), will never have a conclusive or ‘closed’ description. Admittedly, underdetermination, in an epistemological perspective, is not a new notion to philosophy of science. I would like to emphasize, however, that, applied to chemistry, underdetermination has a more ontological connotation to it: The material world is changed when a new substance enters the stage, and the prediction of its actual input is severely restricted. Joachim Schummer discusses three arguments for the creator of a substance to justify his or her actions in a moral discourse: the necessity argument, the knowledge argument, and the
intention argument (Schummer 2001). With respect to the knowledge argument, which is the most pertinent for the present issue, Schummer claims:

[…]every substance bears an infinite potential of properties. From that it follows, however, that it is very likely that any new substances can be used to cause harm. Thus, we may expect that our chemist, while being unable to foresee the particular case of harm, knows well about the high probability of possible harm. Therefore, the knowledge argument turns to the contrary and does not help to excuse our chemist. [Schummer 2001, p. 112]

Typically, non-knowledge outmatches knowledge with respect to new chemical entities, which is a good reason to be careful in whatever application. Hence, the very issue of underdetermination – the knowledge about predominant non-knowledge in chemistry – distinguishes the Contergan case principally from other accidents as, for instance, the Titanic disaster. The only feasible way to eliminate risk would be to stop producing and distributing new substances.

3.3 Who takes responsibility for what?

Because some of the classical schools of philosophical ethics, in their search (or fight) for the last fundament of reasoning, have lost contact with specific practical problems of modern life, particularly in medical and health care, the protagonists of the so-called ‘principlism’ drew up a list of major moral principles which are acceptable to a large group of professionals in the biomedical realm (e.g., Beauchamp & Childress 2009, Wiesing 2012). These principles can be likened to religious commandments, but do not explicitly refer to any fundamental commitment, and are applied without prioritization or hierarchy. Because that is a modern standard approach, which is applied in many ethic committees, and in order to get an appropriate framework for the pertinent discussion, I assume that the main players of the story investigated in the present study are addressable by these principles (which I simplify for the time being). I try to locate these players (substance creators, substance testers, substance distributors, legislators, regulators, consumers) within that framework with respect to the present case study. Here are the four most customary principles:

(i) **Respect for autonomy**: The medical expert or healthcare professional should respect the decision-making capacities and wishes of the patient. He or she should actively support the enabling of individuals to make informed choices. The informed consent is perhaps the most important expression for this respect of autonomy.

(ii) **Non maleficence**: The medical expert or healthcare professional must avoid causing harm. Because all treatments include risks this principle conflicts with (iii).
(iii) Beneficence: This considers the balancing of benefits of treatment against the risks and costs; the healthcare professional should act in a way that benefits the patient/consumer. It is necessary to actively support the benefits for the patient/consumer.

(iv) Principle of justice: The distribution of benefits, risks, and costs must be fair. This principle is more complicated to interpret. (How do we define justice?)

I have shown above that informed consent has been violated severely in the Contergan case. The groups of creators (chemists, pharmacists), testers (pharmacologists, physicians), and distributors (traders, apothecaries, physicians) have misused their customers (consumers and their babies) as guinea pigs or ‘canary birds in the mine of our future’ in blind, uninformed clinical trials outside the clinics. Indeed, when it is not known whether a substance being produced is harmful, then the testers and distributors bear the highest amount of responsibility. Later, the producers carry responsibility, too. The legislators must implement and monitor the drug safety framework, which was, as we have seen, quite miserable in most countries during the 1950s, although these vary in detail. I have to stress here that Frances Kelsey was one of the few persons working in administration who took the regulating principles serious, particularly principle (ii), and was very successful with that.

As to the threat for babies, the situation is different. Babies cannot be recipients of any information, therefore principle (i) becomes a respect for future autonomy, and with respect to that, the mother changes into the role of a responsible actor.

Up to this point I have focused on the role of the creators and distributors of thalidomide and their responsibilities. To gain a more adequate and all-encompassing picture for the ethical discourse, however, it seems necessary to analyze the consumers, too. Between 1957 and 1961, about 5 million people took 300 million daily doses (100 mg). In most of the scientific and public discussions, new-borns, stillborns, and mothers are solely presented as victims without freedom of action. Of course, for the children this is correct, but is it correct for the mothers, too? Is it necessary to take sleeping pills on a regular basis, perhaps over the whole period of pregnancy? Pregnant consumers take responsibility for the health and welfare of their offspring, independent of the ethical conviction or belief (see principles (ii) and (iii)). There are good arguments not to consider insomnia and nausea as illnesses during the first trimester of pregnancy, and if both are not illnesses, then no medication is needed at all. After all, in present-day pregnancy pharmacotherapy, sleep-inducing drugs are no longer used, and thalidomide (and its relatives) is prescribed only for very particular indications (for example, leprosy, cancer, AIDS) and with detailed safety instructions (Schaefer & Weber-Schoen-
dorfer 2009). Frequently, the general advice is not to prescribe or take new and premature drugs (which is in accordance with the lessons we learned from the Contergan disaster). Hence, we should not forget to take into account the role and responsibility of the drug consumers.

Referring to principles (ii) and (iii): To do no harm to the patient/consumer is an old principle of the medical ethos (*primum nil nocere*), and to do good seems to be almost trivial as a medical principle. Nevertheless, with the present case it is useful to recognize that the former has to do with omitting, the latter with active doing. Ironically, the handling of thalidomide turned those principles upside-down: What was omitted and disregarded by the testers and distributors was particularly principle (ii), and the benefits of some enforced sleep (plus the financial benefits of the companies) could by no means compensate for the malformations of new-borns and stillborns.

The interpretation of the principle of doing justice (iv) goes into a similar direction as the points proposed above. Benefits and risks were not distributed fairly.

### 3.4 Beyond anthropocentrism

Another important field of ethical dispute addressing pharmacology and toxicology in general and the Contergan case in particular is the field of animal ethics, which I mention here only briefly. After 1961, many countries tightened their regulations regarding the approval of remedies. In the European Community, for example, full toxicology studies, including teratogenicity tests, have had to be conducted on at least two species of mammals since 1975. These species are specified, referring to the teratogenicity tests, as rabbits on the one hand (of a race that is ‘sensitive against a proven embryotoxic substance’) and rats or mice on the other. Without doubt, this specification is the result of the Contergan case. But who can know in advance whether or not any teratogen reacts like thalidomide? And who can be sure that the chosen animals are of the right species for any risk impact – imaginable or not? The answer to both questions is simple: no one. We still do not know enough about the molecular mechanisms, and it is still possible for a substance to pass the now ‘tightened’ toxicological controls, and nevertheless cause harm on a different path than expected. What certainly has been tightened is the precarious situation of test animals, which have been sacrificed in significantly increasing numbers since Contergan. The main legitimization for this procedure is the fact that there was no comparable case ever since (but this might lead to a typical induction fallacy). Hence, aside from the general use and consumption of animals as substance testers, the Angst-
driven decisions of the legislators and regulators around the world are questionable in both methodological and ethical perspectives.

4. Conclusions

(1) According to Joachim Schummer, those who create and prepare substances carry responsibility for the impact of these substances, and should be aware of providing morally relevant arguments which can support their actions (Schummer 2001, p. 111). Particularly if adverse effects have become known in the case of a remedy, further production becomes questionable from an ethical point of view. In the present context, however, it is not easy to identify and address the very chemist or chemists who prepared the substance for the first time. Along with other disinformational statements, the officials of Grünenthal have not provided trustworthy references about the possible prehistory and the early history of this bioactive substance until the present day. What we have are good arguments that the embryopathic property is not the only pharmaco- and toxicodynamical difference between humans and (most) rodents. Indeed, the question – which is of ethical interest as well – remains: Where did the information about the sleep-inducing impact on humans come from?

(2) As with other pertinent issues of applied ethics – see for example the discourse on active and passive euthanasia – the thalidomide story is foremost about doing and omitting: The main omission of the leaders of Grünenthal – one that is morally inexcusable – has been the continued lack of reports on side effects (as well as other alarming information resulting from the company’s studies). Their equally morally wrong action was the misrepresentation of the facts to the scientific community, physician, and the public. Their behavior described in the present paper was without doubt a scandal.

(3) The embryotoxic effect was a new and unpredictable chemical property of thalidomide; this part of the story was a real accident and no fault (like, for example, the catalytic depletion of stratospheric ozone by fluorochlorohydrocarbons). The characterization of a (new) substance is never finished and sometimes mercurial (cf. DDT, cholesterol, laughing gas, fluorochlorohydrocarbons, and many others).

(4) The dream of an entirely harmless but fully effective remedy should never be dreamt again. To paraphrase the old wisdom of Paracelsus: If it is active, then it has side-effects, too.

(5) There has rarely been an artificial substance that came so close to humans with such dangerous, unexpected, and at the same time promising po-
tential as thalidomide. However, it is not the substance that should be blamed, but those humans who do things by halves.

I shall finish by briefly addressing an issue not yet referred to in the present study, and seldom elsewhere in the body of work dealing with thalidomide. That final issue touches on a different aspect of ethical discourse. It is quite common for many institutions and for the public to project disaster or catastrophe on those humans born with the described symptoms, and these humans are considered to be ‘Sorgenkinder’ (problem children). But does it go without saying that all thalidomiders lead a miserable life and that they suffer on a daily basis? There are several autobiographical accounts, most of which impressively show that this is not the case (Eistel 2007). Bettina Eistel, to mention only one of these thalidomiders, studied psychology, is a very successful dressage equestrienne with medals from European championships and the Paralympics, and is a well-known television moderator. There is much happiness here and creative power, and there is no need to reduce the lives of these persons to their disabilities. Hence, if there is a problem, it might as well be one of the societal environment rather than that of the thalidomiders themselves.

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Notes

1 The motto is taken from an advertisement of the German remedies Contergan and Contergan forte by the company Grünenthal, reproduced in Kirk 1999, p. 267, my translation. ‘Panta rhei’ means ‘all is in flux’, which is ascribed to the Presocratic philosopher Heraclitus.

2 ‘Thalidomide’ is an artificial name, derived from the older abbreviated name α-Pthalimidoglutarimide; IUPAC calls it 2-(2,6-dioxo-3-piperidyl)isoindoline-1,3-dione. The Chemical Abstract System (CAS) number is 50-35-1.
Though usually very rare (Pschyrembel 1986, p. 389), phocomelic malformations of new-borns have not been excluded from public knowledge, see the 'black painting' named 'Mother showing her deformed child to two women' by Goya (reproduced, for example, in Hoffmann 1995, p. 133, and in Roth 2005, p. 212).

The most typical of these malformations of legs and arms are subsumed under 'phocomelia', which refers to the flippers of seals; the more general medical term for malformations of extremities, however, is 'dysmelia'. There are several other aspects of the impact of thalidomide on new-borns.

Teratology is the science of the phenomena, causation, and mechanisms of morphological and functional developmental disorders in animate beings, see Mutschler & Lemmer 1985. Teratogenicity, carcinogenicity, and mutagenicity are the three main classes of chronic toxicity, contrasting with acute toxicity.

Sjöström & Nilsson 1972 is the most original of the thalidomide monographs, that is, all later publications refer to or are based upon the latter. Both authors were involved in the Astra case in Sweden, thus the book is slanted to a certain degree (and it lacks an index). Knightley 1979 is a brilliantly written must-read with an index, and a very informative appendix on the chemistry of thalidomide (information which is almost entirely missing in the other books mentioned here). Kirk 1999 is a doctoral thesis in the history of pharmacy from the University Greifswald, Germany. It commendably meets the requirements for written scientific work, particularly neutrality, although an index is missing. Stephens & Brynner 2001 has a very useful index, but is not throughout reliable referring to details, and perhaps too speculative in some spots. Zichner et al. 2005 does not present material that goes beyond that of the earlier works, particularly that of Kirk. It has its strength in a more sociological and ethical scope, and the integration of orthopaedic aspects (which I do not refer to here). The historical dissertation of Niklas Lenhard-Schramm (2016) is an easy accessible, fresh and substantial documentary, highly recommendable for German speaking readers.

Cf., for example, Schneider 1985, Talbot & Waller 2004. Apart from Contergan, Grünenthal had an intriguing history of pharmacological failures of its own (cf. Sjöström & Nilsson 1972). As to Vioxx, see Rourke 2006 and the references therein.

In an extensive survey about the many new pharmaceutical developments in the second half of the 1950s, the author compiles the formulas of disubstituted barbiturates, Persedon, Noludar, Doriden, and Contergan in a row (Kunz 1959). (It is not clear if this W. Kunz from 'Dr. Schwarz Arzneimittelfabrik GmbH, Monheim' is the same who worked with Grünenthal a few years before.)

'Initial excitation phase' means the higher activity of the animals directly after administration, an effect well-known, for example, for barbiturates (presumably ‘+++’ means very high and ‘−−’ none).

Initially, the following four studies sponsored by Grünenthal have been published in scientific journals: Esser & Heinzler 1956, Jung 1956, Stärk 1956, and Walkenhorst 1957. Later, a study on nursing (not pregnant) women followed (Blasiu 1958). None of these went deeper into metabolism and pharmacokinetics.

Among the first to become aware of possible neuronal damage caused by thalidomide was the physician Ralf Voss from Düsseldorf. As early as 1959 he contacted the Grünenthal officials, but they dismissed the warning. (See the chapter with the telling title ‘How to Sell a Wonder Drug’ in Knightley 1979.) Later, he published his observations (Voss 1961).
See, for example, Lenz 1961, Lenz & Knapp 1962, and, also from Lenz’s working group, Nowack 1965.

By ‘blind’ clinical study I here mean the administration of a presumably active substance to humans without their notice or information and before safety has been established.

Widukind Lenz and his group found out that the first trimester of pregnancy is the most critical phase for the impact of thalidomide.

For the thalidomide story in the USA, see Sjöström & Nilsson 1972, pp. 112-130; Knightley 1979, pp. 64-86; Stephens & Brynner 2001, pp. 39-59; and Kirk 1999, pp. 191-205.

An informative account of that aspect can be found in Roth 2005.

Whereas pharmacodynamics investigates the impact of incoming substances (‘What does the stuff do to the body?’), pharmacokinetics examines what becomes of the incoming substances (‘What does the body do to the stuff?’). For a full understanding of the nature of a substance (including adjunct ethical issues) in the biomedical field, it is necessary to take into account both aspects.

Stephens et al. (2000) give a more recent commentary on the embryopathy explanation hypotheses.

Stephens & Brynner (2001), from whom I borrowed the expression ‘dark remedy’, tell the more recent part of the history in a fascinating, lively and engaged manner. (Stephens was involved in pertinent scientific work for many years, see, for example, Stephens et al. 2000)

Raza 2002 argues for the strict non-use of thalidomide because of the possible misuse.

At the same spot, however, Hoffmann hints at teratological tests that were performed in the 1950s, but does not refer to the uneasy way to find an adequate ‘animal model’.

‘Fraud and Deceit in Science’ is the subtitle of Betrayers of the Truth (Broad & Wade 1982). In this book, the authors present a list of known or suspected cases of scientific fraud. I have shown here that it would be possible to consider the original thalidomide ‘research’ as a candidate for that list.

Using mainly the files of the Alsdorf trial, Beate Kirk gives a meticulously penetrating account of that part of the story in 1960-1961, which in fact is an almost classical sociopathological tragedy about practical constraints (Sachzwänge) in professional life (Kirk 1999, pp. 66-86). She apparently tends to this interpretation; intriguingly, the results of a study initiated and accepted by Grünenthal in 1957 gave an obvious account of the neurological activity of Contergan – as stimulant (!) for electroencephalograms (Walkenhorst 1957).

For detailed accounts of this story see the previously mentioned references. In the German Democratic Republic, thalidomide was not licensed because there were already enough hypnotic remedies, rather than for safety or ethical reasons. Note that the German edition of Sjöström & Nilsson 1972 was published in the GDR. The foreword by the Berlin pharmacologist F. Jung is a practically undisguised piece of propaganda.

Maio 2005 points out that while it can be immoral not to test medicines on humans sufficiently, it can be immoral to test them on certain groups of humans (here: pregnant women).
See, for example, the accounts of that expression in the introductory textbooks Ladyman 2002, French 2007 and Wiltsche 2013. The customary understanding of underdetermination is the lack of data supporting a certain theory.

In a similar sense, the relation between knowledge and non-knowledge is addressed by the Humean induction problem, too: “When we describe a drug as ‘safe’, therefore, what we should really say is that it is a drug that has not yet been found unsafe.” (Knightley 1979, p. 24)

Annas & Elias (1999) compare these two cases. Referring to the re-introduction of the drug in the USA, they say (p. 100): “Like preventing death at sea, preventing thalidomide-affected births will require not only medical technology but also human alertness.”

In a broader sense, a similar canary-bird-metaphor is used by Murphy & Goldkind 2005.

For a short discussion of that point see Nesse 2012, p. 175.

Note that this principle is explicitly used as an accusing claim against the manufacturers by Voss 1961.

We should reformulate principle (ii) as follows: Leave the patient alone if he or she is well!

In Germany the ‘Aktion Sorgenkind’ (‘Initiative Problem Child’) was a fundraising campaign involving a television show and a lottery that was initially motivated by the Contergan story. It was first introduced in 1964, and over the years about 1 billion euros have been collected to support the disabled. In part due to this initiative, many negative stereotypes about disabilities and personal differences have strongly and verifiably changed in Germany. Intriguingly, the name of the project was changed to ‘Aktion Mensch’ (‘Initiative Human’) in the 1990s.

References


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