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Chapter 9

A Review on the Assessment of the Potential Adverse Health Impacts of Carbamate Pesticides

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1. Introduction

Carbamates are an important class of pesticides used worldwide in public health, among rural and urban settings. Indeed, due to their mode of action and effectiveness, the application of these compounds is one of the best options presently offered for pest control in modern agriculture. They are also used for gardening, and as therapeutic pharmaceuticals for veterinary medicine. Carbamates have been also used in medicine for myasthenia gravis, an autoimmune disease which affects the postsynaptic element of the neuromuscular junction, and as pre-exposure protection in military settings from chemical warfare nerve agents such as Sarin and Tabun. For example carbamates such as physostigmine, and pyridostigmine have been listed as human drugs. However, current environmental concerns of the deleterious health impacts of carbamate pesticides have been increasing. Humans and other non-target species are exposed to residues of these cholinesterase-inhibiting chemicals via nutritional sources (legumes, fruits, contaminated meat, and dairy products), water and/or through environmental/occupational settings due to inappropriate handling. As other pollutants, carbamates may induce deleterious effects on both biotic (micro and macro fauna and flora) and abiotic systems. The adverse effects of several carbamate pesticides include renal, hepatic, neurological, reductive, immune, and metabolic functions in both humans and animals. Furthermore, some of them are classified as endocrine disrupting chemicals [1], and regarded as priority pollutants by the United States Environmental Protection Agency (US EPA) [2].

In this chapter a brief overview of the current knowledge on the carbamates’ mode of action and toxicological aspects is presented. The role of in vivo studies (histological and hematological approaches), epidemiology and interdisciplinary research on assessment of the carbamates’ environmental and potential public health effects is addressed and the major...
contributions are discussed. In addition, this chapter presents the results of some of our laboratory experiments that focus on the evaluation of aminocarb and thiodicarb renal and hepatic toxicity, lymphoid organ damage including the spleen and thymus, and adverse effects on male reproductive organs.

2. Mode of action and toxicology

Several key issues on carbamate pesticides such as mechanism of action, and toxicological aspects, including adverse human health effects were recently reported by our group [3]. Briefly, this paper also addresses other topics for a better understanding of toxicological effects, namely risk and exposure assessment, biomonitoring, and analytical methods for the detection of these chemicals on foodstuffs and biological fluids or tissues (e.g. blood, serum, urine, breast milk, hair). International legislation was also mentioned in this report.

Carbamates are esters of N-methyl carbamic acid, known as acetylcholinesterase-inhibiting agents (AChE). As with other pesticides like organophosphates, carbamates inhibit the acetylcholine esterase enzyme which catalyzes the hydrolysis of acetylcholine (Ach), a neuromediator agent, which results in Ach increase at a nerve synapse or neuromuscular junction, thereby increasing stimulation of those nerve endings [3]. Carbamates' cholinesterase-inhibiting effect is reversible compared to organophosphates which is irreversible.

The range of the toxicity of carbamates is variable [5]. Several carbamates have slight (LD₅₀ > 200 mg/kg) to highly (LD₅₀ <50 mg/kg) toxic activity in rodents (Table 1). For example thiodicarb (dimethyl N, N'-thiobis (methylimino) carbonyloxy bisethanimido thioate) is a conventional insecticide for controlling cotton bollworm [6]. It is categorized as class II, moderately toxic, by USEPA and World Health Organization (WHO). In addition, several factors including the route, duration, and frequency of exposure, contact with other pollutants, and compromised physiological condition (e.g. hepatic injury) may determine the degree of toxicity [7-8].

Among other pesticides, some carbamates were included in the list of endocrine-disrupting chemicals (EDCs) by WHO [1]. Due to the potential dangerous effects on both wildlife and human health, this issue has received considerable attention within the scope of public health. The potential of EDCs to interfere with the synthesis, secretion, transport, metabolism, and elimination of a wide range of hormones was already established. These hormonally active chemicals may induce a wide range of deleterious health effects such as developmental, behavioral and reproductive deficits. Agonistic and antagonist mechanism was described. Recently, De Coster and van Larebeke [9] presented an overview of relevant chemicals with endocrine disrupting features, including carbamate pesticides such as chlorpropham, carbaryl, benomyl, methiocarb, pirimicarb, and propamocarb. In this well designed review, authors provided different mechanisms such as the activation of the classical ERα and ERβ nuclear receptors, through estrogen associated receptors, and membrane-bound estrogen-receptors, among others.
3. Potential public health risks

A growing body of literature evidences the harmful effects of carbamates and other pesticides on human and environmental health. Carbamates have been analyzed in environmental analysis, food safety, toxicology, and occupational health. Due to the extensive use of carbamates for agricultural and non-agricultural purposes, their residues have been detected in soils, wastewater effluents, surface water and raw drinking water sources, as well as food products, around the world, and have received particular attention because of their toxicity. The control of the levels of the residues of these compounds in the environment and in crops has an outstanding importance. The presence of pesticide residues and/or their degradation products, which sometimes are more toxic than their precursors in the environment and in foodstuffs, calls for the use of very sensitive analytical methods, capable of determining these compounds at concentration levels equal to or lower than the maximum residue levels (MRLs) established by international organizations [10]. Except for occupational exposure or at home application, e.g. home gardens or the handling with domestic animals, people are exposed to pesticides mainly through diet. Human intake due to pesticide residues in food commodities is usually much higher than those related to water consumption and air inhalation. The evaluation of pesticide residues in food is nowadays a priority objective to ensure food quality and safety, as well as to protect consumers against potential health risks [3, 11]. Considering the chronic exposure through food, Jensen and co-workers [12] used the probabilistic approach to estimate the cumulative exposure to organophosphorus and carbamate pesticide residues from the consumption of fruit, vegetables and cereals in the population of Denmark. Despite the limitations and the uncertainties in the calculation of the dietary cumulative intake, the results showed that exposure for children aged up to 6 years were, on average, 2.4 times higher than the exposure for the general population. Tomatoes were the food source that provided about 67% of the total intake of acetylcholinesterase enzyme-inhibiting pesticides. An outbreak of food borne-illness was reported due to severe methomyl intoxication in Korea [8]. Six elderly people collapsed abruptly after eating 1-2 spoons of boiled rice mixed with bean sprouts and seasoned with soybean sauce. One patient died of cardiac arrest. Symptoms of toxicity presented quickly in the subjects and progressed rapidly, including chest tightness, an unusual sensation in the pit of the stomach, dizziness, ataxia, and finally, collapse. Three patients who drank ethanol with the meal experienced only mild toxic symptoms [8].

The relationships between possible exposure to pesticides and the implications for human health have been the matter of exhaustive and multiple reviews [9, 13-21]. The overall conclusions are that pesticides may induce chronic health complications leading to several diseases. For example neurodevelopmental or behavioral problems, birth defects, asthma, and cancer were documented in children [22].

Due to the relevance of hormones through the life cycle, EDCs may interfere with the developmental processes of humans and wildlife species. The extent of exposure to EDCs may severely affect the most vulnerable life stages including prenatal, early postnatal life, and children. Developmental exposures may induce alterations that, while not evident as birth
defects, can promote permanent changes that lead to increased incidence of diseases throughout life [1].

Several recent reports indicate a correlation between EDCs and numerous chronic diseases such as cancer, diabetes, developmental deficits, obesity, and reproductive health disorders. For example, the potential influence of several endocrine disruptor pesticides on human health was reviewed by Mnif and colleagues [15]. Among other carbamates, aldicarb was demonstrated to inhibit the activity of 17 beta-estradiol and progesterone; carbendazim induced an increase of estrogen production and aromatase activity, although low estrogen effect was reported for carbaryl.

Deleterious health effects of some carbamates chronic exposure in occupational settings were already described, and environmental and public health impacts also considered [14]. Occupational pesticide exposure associated with cancer incidence is thoroughly discussed elsewhere. Alavanja and Bonner reviewed association between carbamates such as aldicarb and carbaryl with colon cancer and melanoma, respectively [23]. In this review and concerning the U.S. Agricultural Health Study (AHS), specific pesticide exposures ascertained by questionnaire prior to the onset of disease were found to be significantly associated with cutaneous melanoma (eg. for more than 56 days of exposure to carbaryl) [23]. Also positive relations between non-Hodgkin lymphoma and carbamate insecticides, among other pesticide exposures in occupational agricultural sceneries, were also lately reported [21]. The relationship between carbamate pesticides, namely carbaryl, and multiple myeloma occurrence was recently described [24].

The association between occupational exposure to organophosphate and carbamate pesticides and semen quality, as well as reproductive and thyroid hormone profiles of Venezuelan farm workers, was undertaken by Miranda-Contreras and colleagues [19]. These findings confirm the potential impact of occupational exposure to EDCs on male reproductive function. Features like sperm chromatin damage and reduction of semen were documented and adverse reproductive health outcomes were detected. The evidence available today shows that both men and women can experience adverse reproductive effects as a result of chronic exposure to carbamate pesticides.

Occupational pesticides non-intentional poisoning was also reviewed recently and carbamates were one of the main groups of pesticides-related mortalities in Brazil [25]. Multiple routes may be considered (inhalation, dermal, oral), with skin contact being one of most common routes of exposure. Protective measures for pesticides exposure related with dermal route of exposure were recently reviewed [26]. In fact, safe handling through personal protective equipment may reduce absorption of those chemicals. In the Republic of Korea, mortality studies due to organophosphate and carbamate poisoned patients (occupationally linked acute exposures or suicides) were recently reported in which 17 cases, under the age of 56.8 ± 19.2 years were found among a total of 146 [7]. Unlike other types of intoxication, there are definite antidotes for carbamates exposure. The mortality of these disease entities could be diminished with sufficient use of atropine, 2-pyridine aldoxime methyl chloride (2-PAM, known as pralidoxime) and vigorous airway management if used from the early stage of their occurrence [7].
4. *In vivo* studies

As for other hazardous chemicals, the contribution of laboratory experimental studies is relevant to understand the impact of carbamate pesticides on public health. Extensive research based on animal experiments, particularly mammals, has evidenced the toxicological effects of a wide range of carbamates. Table 1 displays some relevant contributions of predictive toxicology studies carried out in laboratory animals, based on several approaches. Results clearly show that a number of carbamates have led to a broad spectrum of adverse health effects on different tissues, organs, and systems (hepatic, renal, developmental, and reproductive) in a dose dependent manner with obvious implications on functions. In particular, exposure to carbamates during critical periods of life (eg. pregnancy, and fetal development) induces maternal health anomalies and developmental disability. Generally, apart from hepatic toxicity, harmful effects on reproductive health through altered spermatogenesis, and reduced semen quality have become a noticeable concern.

As shown on Table 1 the most characterized compounds are carbaryl and carbofuran. Alterations on brain, liver, and testis accompanied by testosterone level decay were reported after carbaryl exposure. In addition, nephrotoxic, hepatotoxic and intestinal disruption were documented after intoxication with carbofuran.

Recently, the toxicity of two new carbamates, ethyl-Ś-bromophenyl-carbamate and ethyl-Ś-chlorophenylcarbamate, was characterized [Ŕš, ŘŞ]. Authors reported low subchronic toxicity to rats as evidenced by low severity and reversibility of the majority of the observed alterations [ŔŞ]. Still, degenerative changes in liver, binucleated hepatocytes, and focal coagulative necrosis were noted, and increased lesions were related to high dosage. Biochemical parameters, namely plasma enzymes such as gamma-glutamyltransferase, lactate dehydrogenase, and creatinine exhibited a slight increase.

<table>
<thead>
<tr>
<th>Carbamate Pesticide</th>
<th>Animal</th>
<th>Exposure route &amp; dosing</th>
<th>Tissue/Organ/system</th>
<th>Results</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aminocarb</td>
<td>Rat</td>
<td>Orally, 10, 20 and 40 mg/kg bw for 14 days</td>
<td>Blood, liver and kidney</td>
<td>Hemorrhagic focus on hepatic and renal parenchyma, toxic effects on lymphoid organs</td>
<td>[29]</td>
</tr>
<tr>
<td>Bendiocarb</td>
<td>Rabbit</td>
<td>Orally 5 mg/kg bw daily for 10 and 30 days</td>
<td>Testis</td>
<td>Decrease of testicular weight, degenerative changes on testicular parenchyma, and Leydig cells</td>
<td>[30]</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Per os daily at a dose of 5 mg/kg/bw, and after day 11 received the same dose every 48 h.</td>
<td>Liver</td>
<td>Affects the liver ultrastructure; regeneration of the damaged tissue</td>
<td>[31]</td>
</tr>
<tr>
<td>Carbamate Pesticide</td>
<td>Animal</td>
<td>Exposure route &amp; dosing</td>
<td>Tissue/Organ/system</td>
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<tr>
<td>Ethyl-4-bromphenyl-carbamate and ethyl-4-chlorphenyl-carbamate</td>
<td>Rat</td>
<td>Subchronic oral toxicities; drinking water (12.5, 25 and 50mg/kg/day) for 90 days; 5, 50, 300 and 2000mg/kg single dose using an intragastric tube</td>
<td>Many organs: Lung, brain, cerebellum, intestine, stomach, liver, kidney, heart, and muscle.</td>
<td>Low subchronic toxicity and reversibility on liver, and spleen. Both carbamates are low hazard; signs of toxicity at the higher dosages. The maximum dose of each carbamate did not cause clinical manifestations or liver and skin alterations</td>
<td>[28] [27]</td>
</tr>
<tr>
<td>Carbaryl</td>
<td>Rat</td>
<td>Orally Chlorpyrifos, carbaryl and a mixture for 90 consecutive days, per os</td>
<td>Liver, kidney, urine</td>
<td>No significant histopathological changes; mitochondrial enzymes were affected</td>
<td>[32]</td>
</tr>
<tr>
<td>Carbaryl</td>
<td>Rat</td>
<td>10, 30 mg/kg via intraperitoneal 35 days of exposure</td>
<td>Blood testis</td>
<td>Decline in the testosterone levels; increase in LH and FSH levels Decrease in number of germ cells</td>
<td>[33]</td>
</tr>
<tr>
<td>Carbaryl</td>
<td>Rat</td>
<td>oral gavage at 2 ml/kg preweaning age to senescence</td>
<td>Brain, plasma, liver</td>
<td>Dose-related increase at all ages, with differences across life span</td>
<td>[34]</td>
</tr>
<tr>
<td>Carbaryl</td>
<td>Rat</td>
<td>Orally at 0, 20, 100 and 200 mg/kg for 80 days prior to mating.</td>
<td>Testis</td>
<td>Adverse effects on spermatogenesis, resulting in reduced fertility</td>
<td>[35]</td>
</tr>
<tr>
<td>Carbendazim</td>
<td>Mouse</td>
<td>Mated mice 0, 150, 300, and 600 mg/kg/day by gavage.</td>
<td>Maternal Blood Fetuses</td>
<td>Dose of 150 mg/kg/day induced a very slight increase in postimplantation loss; maternal and developmental toxicity at 300 and 600 mg/kg/day</td>
<td>[36]</td>
</tr>
<tr>
<td>Carbendazim</td>
<td>Rat</td>
<td>Once daily p.o. at 10 ml/kg for a dose of 200 mg/kg/day.</td>
<td>Testis</td>
<td>Tubular dilation, tubular necrosis, and/or germ cell degeneration</td>
<td>[37]</td>
</tr>
<tr>
<td>Cartap, carbosuran</td>
<td>Rat</td>
<td>Each pesticide per se (50% LD50);</td>
<td>Serum</td>
<td>Alterations in the serum lipid profile; marked decrease in</td>
<td>[38]</td>
</tr>
<tr>
<td>Carbamate Pesticide</td>
<td>Animal</td>
<td>Exposure route &amp; dosing</td>
<td>Tissue/Organ/system</td>
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<td>Reference</td>
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<tr>
<td>Carbofuran</td>
<td>Rat</td>
<td>4.0 mg/kg/bw for 7 days or 2.8 mg/kg/bw for 30 days daily by Ryle’s tube.</td>
<td>Small Intestine</td>
<td>Intestinal disruption of the villi, and comet assay showed disintegration of DNA in enterocytes of animals exposed for 30 days; toxicity may modulate digestive functions in intestine.</td>
<td>[39]</td>
</tr>
<tr>
<td>Carbosulfan</td>
<td>Rat</td>
<td>Orally 1 mg/kg/bw dissolved in sunflower oil daily for 28 days</td>
<td>Kidneys blood</td>
<td>Nephrotoxic effects through augmented oxidative stress and attenuated antioxidant defense system.</td>
<td>[40]</td>
</tr>
<tr>
<td>Methiocarb</td>
<td>Rat</td>
<td>P.o at 0.5–5 mg/kg/bw for 5 weeks</td>
<td>Liver, bone marrow</td>
<td>Liver toxicity and clastogenic effects (micronucleated polychromatic erythrocytes).</td>
<td>[41]</td>
</tr>
<tr>
<td>Methomyl</td>
<td>Mouse</td>
<td>Orally at doses 25, 10, and 2 mg/kg/bw for 1, 5, and 28 days</td>
<td>Liver, kidney, brain, and testis</td>
<td>Possible lipid peroxidation, disturbances on the GSH levels in liver, kidney, testis, and brain.</td>
<td>[43]</td>
</tr>
<tr>
<td></td>
<td></td>
<td>25 mg/kg/bw for 20 days, (i.g.)</td>
<td>Liver, and kidney</td>
<td>Oxidative damage on liver and kidney, which were partly, ameliorated by the pretreatment of vitamin E and taurine.</td>
<td>[44]</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Orally; 1 mg; 2 mg, 3 mg and 4 mg/kg/bw for 30 days and effective dose of 4 mg/kg/bw for 5, 10, and 20 days</td>
<td>Liver and serum</td>
<td>Harmful effects on cell metabolism, cell membrane permeability, and hepatic detoxification system.</td>
<td>[45]</td>
</tr>
</tbody>
</table>
Table 1. Laboratory animal findings on main carbamate pesticides effects. Abbreviations: bw-body weight; i.g.- intragastrically.

<table>
<thead>
<tr>
<th>Carbamate Pesticide</th>
<th>Animal</th>
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<th>Tissue/Organ /system</th>
<th>Results</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbaryl</td>
<td>Rat</td>
<td>Orally daily for 65 days at 2 doses (0.5 and 1.0 mg kg⁻¹ bw)</td>
<td>Testis, epididymis, and serum</td>
<td>Decreased the fertility index, testicular damage, sperm quality affected</td>
<td>[46]</td>
</tr>
<tr>
<td>Pirimicarb</td>
<td>Mouse</td>
<td>Oral gavage 2.14, and 10.7 mg/kg/day pirimicarb, and dichlorvos plus pirimicarb daily for 30 consecutive days</td>
<td>Liver and serum</td>
<td>Prominent changes in liver oxidative markers, as endogenous metabolites in serum and liver; PI, either alone or in combination lead to changes on liver glucose, fat and protein metabolism, energy metabolism and oxidative balance</td>
<td>[47]</td>
</tr>
<tr>
<td>Thiodicarb</td>
<td>Rat</td>
<td>Intraperitoneal, 2.9 and 5.8 mg/kg daily for 28 days</td>
<td>Vital organs, particularly liver and heart</td>
<td>Significant increase in AST on 7th day. No much changes on the various biochemical profiles except inhibiting AChE. No adverse specific damage to vital organs, mainly liver and heart</td>
<td>[49]</td>
</tr>
</tbody>
</table>

4.1. Case studies: Effects of thiodicarb and aminocarb on male reproductive system

As presented in Table 1, the effects of thiodicarb on various biochemical parameters and blood enzymes were investigated in adult male wistar rats following its daily intraperitoneal administration at rates of 2.9 and 5.8 mg/kg for 28 days [49]. The findings of this research indicated that thiodicarb did not significantly affect or alter the various biochemical profiles except inhibiting AChE following intraperitoneal administration up to 28 days.

The systemic toxic effects of thiodicarb on rats have been well described by our group [48]. Several approaches such as hematological, biochemical, histopathological, and flow cytometry were used in this paper to characterize the subacute effects. Marked systemic organ toxicity was reported including renal and testis degeneration, appreciated cellular loss on thymus, hemorrhagic focus on liver, and disruption within the spleen. T lymphocytes displayed high values. This paper also evidenced some hemorrhage on interstitial tissue of testis.

To complement the above mentioned work, and in order to fully characterize the effects of thiodicarb on the male reproductive system of rats, experiments were conducted under the
guidelines for ethics on animal experimentation using a similar protocol for epididymis (another reproductive organ). Briefly, three months old rats purchased from Harlan Iberica (Spain) were divided into two groups, and kept under appropriate conditions. Thiodicarb was dissolved in water and was given every day (40 mg/kg body weight) for a period of 30 days. For comparison, animals given water only were used. After one month, animals were anesthetized, and sacrificed for epididymis sampling and further histological analyses. No apparent macroscopic changes were noted in organs of thiodicarb-exposed rats. However the observations at light microscope level evidenced a reduction of sperm mass within the lumen (Figure 1). However no changes were noted on the epithelium lining the ducts.

Figure 1. Histological sections of rat epididymis (E) from (a) control; and thiodicarb (40 mg/kg body weight) exposed animals during one month period (b). A considerable decrease on sperm is noted in the lumen (arrow); Inset – a general view displaying a reduction on sperm (arrow); haematoxylin and eosin stain. Original magnification: (a) x200; (b) x200; inset x40.

Aminocarb (4-dimethylamino-3-methy-N-carbamate) is a phenylsubstituted methylcarbamate pesticide broadly used to control the growth of insect pests such as Lepidoptera and Coleoptera species affecting agriculture and storage of legumes, fruits, and grains. The toxicity of this carbamate on rats was thoroughly characterized through histological, hematological, and biochemical approaches as shown on Table 1. The results of this study evidenced multi-organ damage and the extension of lesions were dose-dependent. Studies on progress using a similar experimental dosing procedure (30, and 40 mg/kg body weight, respectively) were conducted in male rats aiming to evaluate the effects of aminocarb on testis and epididymis using histological assays. The results clearly show harmful effects on both testis and epididymis (Figures 2 and 3). Epididymis revealed a reduction of sperm compared to control. The testis presented some degenerative changes such as vacuolation, germ cell loss with obvious decrease of seminiferous epithelium, and release of immature germ cells into the lumen.
Figure 2. Epididymis (E) of (a) control and aminocarb-dosed rats (20mg/kg) for 14 days (b) displaying a decrease on sperm (arrow); testis (T) from control (c) and pesticide exposed rat (d-e) evidencing strong vacuolation (*), and immature germ cells (arrow); haematoxylin and eosin stain. Original magnification: (a,c,e) x$100$; (b) x$200$; (c-e) x$200$.

Figure 3. Representative histological section of testis from aminocarb-exposed rat (40 mg/kg/body weight) during 14 days. Immature germ cells (arrow) within the lumen of seminiferous tubules; some tubules (*) denoted a decrease on germ cell layers; haematoxylin-eosin stain. Original magnification: x100.
Overall taken together, the results mentioned above clearly evidence the deleterious effects of both carbamates (thiodicarb and aminocarb) on male organs namely testis and epididymis. In fact disrupted spermatogenesis, and subsequent changes in epididymis ducts may compromise the reproductive potential. These findings are consistent with the results from studies on other carbamates reported on Table 1.

5. Conclusion

Although efforts have been made globally and a significant progress was accomplished, the impact of carbamates (mainly of those that exhibit endocrine disruptor behavior) on human and environmental health still remains a public health problem and a challenge. Insights from endocrine disruptor research in animals have a huge impact on current practice in toxicological evaluation. The effects of exposures should be studied in adulthood but also, and particularly, in fetal development, perinatal life, childhood and puberty.

Continuous efforts to undertake multidisciplinary research based on in vitro technologies and in vivo toxicological studies, coupled to the epidemiological studies of exposure in humans are mandatory in order to improve our knowledge on the underlying mechanisms and health consequences. Also, protection programs, including educational ones, on the appropriate use of pesticides to minimize population exposures as well as preventive health monitoring are needed principally in developing countries.

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