Endocrine-disrupting activity of mancozeb

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Abstract

The objective of the present study was to review the existing data on the mechanisms involved in the endocrine disrupting activity of mancozeb (MCZ) in its main targets, including thyroid and gonads, as well as other endocrine tissues that may be potentially affected by MCZ. MCZ exposure was shown to interfere with thyroid functioning through impairment of thyroid hormone synthesis due to inhibition of sodium-iodine symporter (NIS) and thyroid peroxidase (TPO) activity, as well as thyroglobulin expression. Direct thyrotoxic effect may also contribute to thyroid pathology upon MCZ exposure. Gonadal effects of MCZ involve inhibition of sex steroid synthesis due to inhibition of P450scc (CYP11A1), as well as 3β-HSD and 17β-HSD. In parallel with altered hormone synthesis, MCZ was shown to down-regulate androgen and estrogen receptor signaling. Taken together, these gonad-specific effects result in development of both male and female reproductive dysfunction. In parallel with clearly estimated targets for MCZ endocrine disturbing activity, namely thyroid and gonads, other endocrine tissues may be also involved. Specifically, the fungicide was shown to affect cortisol synthesis that may be mediated by modulation of CYP11B1 activity. Moreover, MCZ exposure was shown to interfere with PPARγ signaling, being a key regulator of adipogenesis. The existing data also propose that endocrine-disrupting effects of MCZ exposure may be mediated by modulation of hypothalamus-pituitary-target axis. It is proposed that MCZ neurotoxicity may at least partially affect central mechanisms of endocrine system functioning. However, further studies are required to unravel the mechanisms of MCZ endocrine disrupting activity and overall toxicity.

Keywords: mancozeb, thyroid, testes, ovary, adrenal, adipose tissue

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Introduction

Mancozeb (MCZ) is a dithiocarbamate fungicide widely used since 1962 for crop protection against a wide spectrum of fungal pathogens (1). Close relatives of MCZ from the same group of fungicides include Maneb and Zineb containing Mn$^{2+}$ and Zn$^{2+}$ ions, respectively (Figure 1).

Environmental persistence of MCZ in soils was considered rather low (2), with half-life from 2 to 8 days in aerobic and anaerobic soils, respectively (3). However, MCZ-affected soils were characterized by an increase in the formation of ethylthiourea (ETU), the main MCZ metabolite, for 30 days with a subsequent decline (4). Therefore, hazardous effects of MCZ exposure are expected to be observed at frequent use. At the same time, application of MCZ was shown to result in 20-fold increase in soil toxicity, with toxicity observed within more than 0.5 km from the application site (5). Although MCZ does not accumulate in plants, certain foods were shown to contain pesticide residues, thus posing significant health risks (6).

Although health hazards of mancozeb exposure are considered low in the case of safe occupational hygiene procedures (7), its wide use poses significant health hazards (8), especially in agricultural communities in developing countries (9).

It is considered that MCZ toxicity is widely attributed to toxic effects of its metabolite, ethylthiourea (Figure 2). Correspondingly, urinary ETU levels were found to correlate significantly with spraying mancozeb (10). However, certain studies demonstrate that adverse effects of MCZ exposure may be ETU-independent (11).

In human and experimental studies, MCZ was reported to have adverse effects on the immune system (12), gastrointestinal tract (13, 14), cardiovascular system (15), as well as carcinogenic effects (16). An increasing body of data has demonstrated profound reproductive, developmental, and thyroid-related toxicity of MCZ (17-19). Moreover, it has been shown that the thyroid and gonadal endocrine-disrupting effects upon MCZ exposure manifest without overt hepatotoxicity, indicative of endocrine system as a primary target for MCZ adverse effects (20).
Figure 2. Formation of ethylenethiourea, a toxic metabolite of dithiocarbamate fungicides.

Slika 2. Nastajanje etilen tioureje, toksičnog metabolita ditiokarbamatskih fungicida.

Concomitantly, the mechanisms of MCZ toxicity in general, and its adverse effects on the endocrine system in particular, have yet to be fully characterized. Therefore, the objective of the present study was to review the existing data on the mechanisms involved in the endocrine disrupting activity of MCZ in its main targets, including the thyroid and gonads, as well as other endocrine tissues that may be potentially affected.

**Thyroid**

MCZ was shown to accumulate to a significant extent in the thyroid following exposure (16), thus predisposing to a wide spectrum of thyroid-associated pathologies. Particularly, lifetime exposure to MCZ was found to be positively associated with circulating higher T3 levels (17), although another study revealed an association between exposure rates and both hypo- and hyperthyroidism (18). However, no significant association between MCZ exposure and thyroid cancer was observed in farmers using fungicide (19). Examination of MCZ-exposed workers revealed significant increase in iodine excretion in association with exposure rates, indicative of thyroid-disrupting effects of MCZ and its metabolite ethylenethiourea (19).

Experimental studies have also shown a significant effect of MCZ on thyroid functioning, as evidenced by reduced T3 and T4 levels, although increased TSH concentrations in response to low T4 levels may be indicative of an adequate negative feedback mechanism. Reductions in thyroid hormone production were also associated with thyroid gland morphology, characterized by epithelial damage, altered stroma, and impaired follicle shape in birds (25). In pregnant rat dams, MCZ exposure also resulted in a dose-dependent decrease in T4 levels, although offspring T4 levels and thyroid
weight were unaffected (26). However, in another study, maternal MCZ exposure resulted in morphological alterations in the thyroid, characterized by a high number of columnar vacuolated cells, indicative of higher secretory activity of the gland in rats (27). Taken together, these findings are indicative of altered thyroid hormone secretion upon MCZ exposure.

The underlying mechanisms of impaired thyroid functioning may involve the interference of MCZ with thyroid hormone synthesis, including inhibition of thyroid peroxidase (TPO) activity \textit{in vivo} (28), which was also supported by molecular docking analysis (29). The inhibition of TPO would be expected to result in altered oxidation rate of iodine and its organification. In addition, MCZ was shown to compete with T3 for binding thyroid hormone receptor (30). Sodium-iodide transporter (NIS) may be also considered as the potential target for MCZ (31), thus reducing iodine availability for thyrocytes and thyroid hormone synthesis.

Other studies demonstrate that the MCZ metabolite ETU is responsible for its thyroid-disrupting activity. Specifically, comparative analysis demonstrated that both mancozeb and especially ETU were considered as thyroid disruptors, reducing intrafollicular T4-content in zebrafish eleutheroembryos by 50% and nearly to zero values, respectively (32).

ETU is known to be a potent TPO inhibitor with IC50 of 0.791 μM (33), thus affecting a key step in thyroid hormone biosynthesis. Despite indications on the potential impact of ETU on NIS activity (34), a recent study failed to corroborate a selective inhibition of the transporter by ETU (31).

ETU-induced thyroid hormone imbalance may be also mediated by its direct thyrotoxic effects. In particular, \textit{in vitro} transcriptomic analysis and \textit{in vivo} study in ETU-exposed mice demonstrated significant thyrotoxicity of ETU that was associated with a significant reduction in thyroid thyroglobulin and Bcl2 mRNA expression levels, altogether resulting in a decrease in free T4 levels (35).
Despite its thyroid-targeted effects, the most recent analysis failed to show a significant relationship between ETU exposure rates and thyroid cancer, contrary to the reported carcinogenic effect of MCZ (36). However, in banana plantation workers, blood ETU levels were found to be significantly associated with thyroid nodule size (37).

**Testes**

Experimental studies demonstrated a significant association between mancozeb exposure and reduced testosterone levels, due to toxic impact on testes and especially testosterone-producing Leydig cells, as well as mechanisms of testosterone biosynthesis. Specifically, oral exposure to MCZ also resulted in a significant alteration of testicular morphology, with Sertoli cells (38) and Leydig cells (39) degeneration in rodents being the most prevalent. Moreover, a significant reduction in testes size, seminiferous tubule diameter, and germ cell and Sertoli cell number was observed in mice exposed to MCZ in utero and during lactation (40), indicative of prenatal endocrine-disrupting effect of MCZ on the male reproductive system.

MCZ exposure in male mice resulted in a significant decrease in basal plasma testosterone levels. Leydig cells isolated from these animals were also characterized by
reduced basal testosterone and pregnenolone secretion, which may occur secondary to
down-regulation of CYP11A1 activity (41). In addition, inhibition of testicular 3β-
hydroxysteroid dehydrogenase and 17β-hydroxysteroid dehydrogenase is considered to
be the potential mechanism for MCZ (500 mg/kg b.w)-induced testosterone reduction
(42). However, another study demonstrated an increase in testicular 3β-HSD activity in
response to 100 mg/kg b.w MCZ exposure (43). Such an inconsistency may arise from
distinct effects of various doses of MCZ, which should be addressed in future dose-
response studies. Moreover, MCZ-induced alteration 3β-HSD activity was accompanied
by the disruption of testicular germinal epithelium, reduction of spermatogenic cells, and
vacuolization of Leydig cells, as well as a significant decrease in serum FSH, LH, and
testosterone concentrations in rabbits. Supplementation with GSH was shown to reverse
these changes, implicating a role of oxidative stress in testicular MCZ-induced toxicity
(43). In turn, apoptotic cell death was shown to follow MCZ-induced oxidative stress in
testes (44).

In addition to reduced testosterone concentrations and impaired spermatogenesis,
MCZ exposure also significantly decreased circulating estradiol and progesterone levels
(15).

In parallel with reduced testosterone synthesis, MCZ was shown to affect its
signaling through a down-regulating effect on the androgen receptor, as well as a dose-
dependent inhibitory effect on agonist-induced AR transactivation (45). Correspondingly,
MCZ possessed anti-androgenic activity in NIH3T3-AR-Luc cells treated with
testosterone (46).

Taken together, the existing data demonstrate that MCZ may alter testosterone
production via direct cytotoxicity to testosterone-producing Leydig cells, as well as
interference with testosterone synthesis and signaling. In addition, reduction of LH in
parallel with MCZ-induced decrease in testosterone production may be indicative of
disruption of the negative feedback mechanism of the hypothalamus-pituitary-gonadal
axis. At the same time, high-quality epidemiological data linking MCZ exposure to male
infertility in humans seem to be lacking.

**Ovaries**

The ovaries are considered to be one of the primary targets for MCZ-induced
docrine disruption. At the organ level, MCZ exposure has been shown to cause a
significant decrease in protein and glycogen content in ovaries, whereas the level of total
lipids, phospholipids, and neutral lipids was increased in parallel with a reduction in
ovarian size and corpus luteum number in rats (47). These changes were also associated
with a significant decrease in follicle number and the number of cycles, whereas the
number of atretic follicles increased substantially in response to MCZ exposure in mice
(48). Similar patterns were characteristic of MCZ-induced inhibition of compensatory
ovarian hypertrophy in hemicastrated female rats (49).
MCZ-induced decrease in follicle number was shown to be associated with apoptosis that was ameliorated by vitamin C and E supplementation, implicating oxidative stress in MCZ-induced ovarian toxicity in mice (50). A combination of these vitamins was also shown to be beneficial in the improvement of oocyte maturation, fertilization and implantation rates, and embryo development in first-generation mice pups from MCZ-exposed dams (51).

MCZ induced a dose-dependent cytotoxic effect on granulosa cells, characterized by altered intracellular contacts, membrane damage, chromatin condensation, cytoplasmic vacuolization, and reduced mitochondria length in mice (52). Given the role of granulosa cells in estradiol synthesis, MCZ cytotoxicity may result in reduced estradiol synthesis. Alteration of adrenal estradiol synthesis may also contribute to female reproductive system dysfunction (53).

A significant decrease in granulosa cell viability in response to mancozeb treatment was associated with increased progesterone levels. Taken together with the observed decrease in oocyte cumulus expansion and maturation upon MCZ exposure, this suggests that increased progesterone production may inhibit ovulation in caprines (54).

It is proposed that the mechanisms underlying mancozeb-induced granulosa cell dysfunction may involve oxidative stress, mitochondrial dysfunction, impaired ATP production, as well as p53 down-regulation (55). The observed decrease in both mRNA and protein p53 expression in MCZ-exposed granulosa cells was associated with increased cell migration, thus inducing premalignant changes (56). In contrast, granulosa cells from MCZ-exposed mice were characterized by a higher apoptotic rate, which could be reversed by antioxidant resveratrol treatment (57), corroborating the previously discussed association between MCZ exposure and oxidative stress in granulosa cells.

In contrast, progesterone synthesis in bovine luteal cells was inhibited by mancozeb exposure, being indicative of MCZ-induced inhibition of P450scc (CYP11A1) and 3beta-HSD (HSD3B1) enzymes (58). Correspondingly, chemical modeling analysis revealed capability of MCZ methyl group to interact with hydroxyl group of Asn315 residue in CYP11A1 (59).

As a result of endocrine dysfunction, exposure of female mice to mancozeb was shown to reduce meiotic maturation of the oocytes and its arrest in metaphase II, as well as decreased formation of oocyte pronuclei following in vitro fertilization (60). Even in oocytes reaching metaphase II, mancozeb exposure resulted in significant alterations of meiotic spindle formation, predominantly characterized by impaired microtubule length (61).

The above-mentioned changes in the ovarian function due to MCZ exposure were shown to be associated with impaired implantation in mice (62). Reduced trophoblastic spheroid attachment to endometrial epithelial cells in MCZ exposed animals may be at least partially mediated by down-regulation of endometrial estrogen receptor β receptor and integrin β3 expression, whereas ETU itself did not possess such an inhibitory effect.
In addition, even in the lack of significant changes of decidualization markers in endometrial stromal cells, MCZ suppressed decidualization-induced morphological changes in stroma cells that may be associated with altered Prostaglandin E synthase transcription and protein expression (64).

In contrast to male effects, the existing epidemiological data demonstrate that MCZ exposure may significantly affect female fertility and adverse pregnancy outcomes, resulting in altered menstrual cycle, spontaneous abortions, miscarriage, as well as adverse developmental outcome in children (17).

Adrenals

Several studies demonstrated that the adrenals may also be considered as potential targets for mancozeb endocrine-disrupting activity. Specifically, even in the absence of cytotoxic effects in rainbow trout adrenocortical cells, MCZ exposure was shown to inhibit dibutyryl-cAMP- and ACTH-induced cortisol secretion (65). Molecular docking analysis of the interaction between MCZ with steroidogenic enzymes demonstrated that MCZ was characterized by the highest binding affinity to CYP11B1, a 11-beta-hydroxylase of the adrenal cortex involved in the biosynthesis of cortisol and corticosterone (66). Moreover, in an angiotensin-II-responsive steroid-producing adrenocortical cell line H295R, mancozeb exposure was shown to be associated with estradiol levels (67).

Adipose tissue

In the last decades, adipose tissue has been considered as a novel endocrine organ secreting a wide spectrum of molecules with systemic effects (68). Certain studies demonstrated the potential impact of MCZ exposure on PPARγ, the key adipogenesis regulator (69). Specifically, in in silico molecular docking analysis, MCZ was shown to bind PPARγ through interaction between thiomethyl and amino acid residues of the fungicide with Arg288, Ser289, His323, and Tyr327 residues in PPARγ molecule (70).

At the same time, in an in vitro study mancozeb was found to be cytotoxic to adipocytes without any significant influence on mPPARγ activation (71). In agreement with this, MCZ exposure was shown to reduce adipogenesis in a culture of adipocytes, although it increased model adipose tissue vascularization in non-toxic concentrations (72).

Neurotoxicity as a potential contributor to mcz endocrine disrupting activity

The existing studies of endocrine disrupting activity of MCZ directly or, more frequently, indirectly implicate the role of altered hypothalamus-hypophysis-gland (thyroid, adrenal, or gonad) in mediating the observed adverse effects. Despite the lack of clear evidence, it has been proposed that toxic effects of MCZ and ETU exposure on neuronal structures may result in endocrine dysfunction due to the role of hypothalamus...
as a central controller of endocrine system functioning (73). This hypothesis was indirectly supported by the role of neurotoxin-induced hypothalamic damage in the dysregulation of hypothalamic-pituitary axis (74). In this regard, MCZ neurotoxicity will be briefly reviewed herein.

Both MCZ and maneb exposure were shown to possess neurotoxic effects in rat dopamine and GABA mesencephalic populations, inducing mitochondrial dysfunction characterized by the inhibition of respiratory chain complex I activity. It is noteworthy that neurotoxic effects were mediated by both organic moiety and Mn2+ (75). Correspondingly, DA and GABAergic neurons were characterized by degeneration in MCZ-exposed C. elegans (76). These findings corroborate earlier indications of Mn2+ toxicity (77) with a special focus on dopaminergic and GABAergic neurons (78). It has been also demonstrated that microglia activation may significantly contribute to MCZ-induced H2O2 generation in mesencephalic dopamine and GABA cell populations, which was also found to be Mn2+, but not Zn2+-dependent (79). Moreover, increased Mn2+ brain accumulation following MCZ exposure was associated with reactive oxygen species (ROS) overproduction and a compensatory activation of Nrf2/ARE signaling in brain (80). Finally, MCZ exposure was shown to increase APP and Aβ42 protein expression in PC12 cells, as well as activate the double-stranded RNA dependent protein kinase that is known to promote neuronal apoptosis (81). In addition, proapoptotic signaling was characterized by modulation of Bcl-2, Bax, Caspase 3, and caspase 9 mRNA expression in adult zebrafish brain (82).

Several studies have also demonstrated the adverse effects of MCZ exposure on neurotransmitter metabolism. Specifically, the levels of excitatory transmitters, glutamate and aspartate, were reduced in the early postnatal period, whereas at the age of one month cerebellar cortex was characterized by the elevation of glutamate and reduction of aspartate (83). An increase in excitatory neurotransmitter release in response to MCZ exposure was also associated with mitochondrial and synaptic dysfunction, as well as neuroinflammation (84). These findings generally corroborate the observation of MCZ-induced inhibition of ATP-dependent vesicular uptake of glutamate (85).

MCZ-induced neurodegeneration at diencephalic sites in Ornate wrasses was also associated with increased orexin mRNA and protein expression, being associated with reduced food intake (86).

It is also notable that behavioral deficits in MCZ-exposed C. elegans may precede profound structural damage in response to MCZ exposure (87), indicating early neuronal effects in the course of MCZ toxicity.

Consistently with the laboratory data, human biomonitoring studies demonstrated that MCZ exposure was shown to affect neurodevelopment in children. Specifically, MCZ exposure as assessed by increased ETU concentrations was shown to be associated with poorer verbal learning ability (88), impaired social-emotional development (89), and semantic memory (90). Although MCZ neurotoxicity may underlie these effects, it is also
proposed that altered sex hormone dysregulation or thyroid dysfunction may also affect the developing brain.

**Conclusions**

The existing data demonstrate that MCZ possesses significant endocrine-disrupting properties in the thyroid and gonads, inducing its effects through toxic damage to hormone-producing cells, inhibition of the enzymes involved in hormone biosynthesis, modulation of hormonal receptors, as well as dysregulation of hypothalamus-pituitary-gland axis. At the same time, an increasing body of data demonstrates that other endocrine tissues, including adipose tissue and adrenals, may be the targets of MCZ toxicity (Figure 4).

**Figure 4.** Targets of mancozeb endocrine-disrupting activity and the potential role of neurotoxicity in endocrine dysfunction through alteration of hypothalamic-pituitary axis.

In addition, due to the role of hypothalamic structures in central control of endocrine system functioning, it has been proposed that MCZ neurotoxicity may at least partially contribute to endocrine disrupting effects of the fungicide. However, further studies are required to unravel the mechanisms of MCZ endocrine-disrupting activity and overall toxicity.

**Conflict of interest**

The authors declare no conflict of interest.
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Ethics statement: N.A

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Delovanje mankozeba kao endokrinog ometača

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Kratak sadržaj

Ovo istraživanje je imalo za cilj da pruži pregled postojećih podataka o mehanizmima uključenim u delovanje mankozeba (MCZ) kao endokrinog ometača u ciljnim organima, u koja se ubrajaju štitna i polne žlezde, kao i u ostalim endokrinim tkivima na koja potencijalno može uticati MCZ. Pokazalo se da izloženost mankozebu remeti rad štitne žlezde tako što ometa sintezu tiroidnih hormona inhibirajući aktivnost natrijum-jodid simportera (NIS) i tiroidne perioksidaze (TPO), kao i ekspresiju tiroglobulina. Neposredno tirotoksično dejstvo takođe može doprineti razvoju tiroidne patologije pri izlaganju mankozebu. Delovanje mankozeba na polne žlezde uključuje inhibiciju sinteze polnih steroida usled inhibicije P450scc (CYP11A1), kao i 3β-HSD i 17β-HSD. Uporedo sa promenama u sintezi hormona, pokazalo se da MCZ snižava slanje signala androgenih i estrogenih receptorima. Ovi efekti koji se odnose na polne žlezde zajedno dovode do pojave reproduktivne disfunkcije kako kod muškaraca, tako i kod žena. Pored jasnih utvrđenih ciljnih organa na koja MCZ deluje kao endokrini ometač, mogu biti pogođena i druga endokrina tkiva. Konkretno, pokazalo se da ovaj fungicid utiče na sintezu kortizola, moguće posredstvom modulacije aktivnosti CYP11B1. Pored toga, utvrđeno je da izloženost mankozebu remeti signale PPARγ, koji su ključni regulator adipogeneze. Postojeći podaci takođe ukazuju na to da do posledica izloženosti mankozebu može doći posredstvom modulacije ose hipotalamus-hipofizaciiljni organ. Navodi se da je moguće da neurotoksičnost mankozeba makar delimično utiče na centralne mehanizme rada endokrinog sistema. Ipak, neophodna su dalja istraživanja kako bi se razjasnili ne samo mehanizmi delovanja mankozeba kao endokrinog ometača već i njegova toksičnost uopšte.

Ključne reči: mankozeb, štitna žlezda, testisi, jajnici, nadbubrežna žlezda, adipozno tkivo