EDTA: Ethylene Diamine Tetra Acetic Acid – A Review

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Abstract

Chelation therapists around the world incorporate chelation therapies into their daily medical practice, frequently using EDTA (ethylene diamine tetra acetic acid) compounds, unaware of the chemical difference of the various EDTA chelating agents. With this information, we aim to clarify the different mode of action of the EDTAs, including their appropriate medical use. In the USA, medical practitioners promote EDTA chelation, often as an alternative to conventional treatments for a variety of chronic diseases, including vascular problems. German nonmedical professionals use the ‘CaEDTA push’ as promoted by US web pages, although this is against standard protocol. CaEDTA has been FDA-approved for lead intoxication only, and NaMgEDTA has not been approved for the treatment of cardiovascular disease. These facts are often overlooked. Misunderstandings increase the risk of iatrogenic accidents. This information aims to prevent this.

Keywords: EDTA; CaEDTA push; Chelation; Lead; Calcium

Na$_2$EDTA, NaMgEDTA and NaCaEDTA Chelation - Understanding the Difference

In the early 1930, the German scientist Munz first synthesized Na$_2$EDTA (disodium ethylene diamine tetra acetate dehydrate or edetate disodium). The chemical was patented and used to soften the hard and calcium-rich waters of Germany, which helped the textile industry to develop more uniform dying processes. In the 1950s, Na$_2$EDTA was FDA (Food and Drug Administration) approved for the treatment of hypercalcemia and digitals intoxication. At about the same time, NaCaEDTA (disodium calcium EDTA), also called edetate calcium disodium, sold as calcium disodium versenate and Hereweth referred to as CaEDTA) was used to treat acute cases of lead intoxication in children and adults [1]. Since then, the FDA has issued the following warning:

“The two EDTA drugs have established names that are easily confused and both are referred to in clinical practice as 'EDTA'. This confusion has resulted in medication errors in which some patients have received the wrong drug, which has been fatal in some cases or caused serious adverse reactions in others. The error is especially dangerous when edetate disodium is erroneously given to a patient who is supposed to receive edetate calcium disodium.”

The two EDTA drugs have different approved uses and significantly different effects. For example, edetate disodium is more likely to cause severe decreases in blood calcium levels. A severe decrease in blood calcium levels due to the erroneous administration of edetate disodium has resulted in death, predominantly among pediatric patients, who were to be treated for lead poisoning with edetate calcium disodium. As noted, FDA has special concerns regarding the use of edetate disodium and is reconsidering the overall risks and benefits of the drug” [2].

Since the 1970s, American medical doctors have added magnesium to Na$_2$EDTA, changing it into NaMgEDTA. Members of The American College of Advancement in Medicine (ACAM) actively promoted the use of NaMgEDTA (dismagnesium EDTA) infusions for the treatment of cardiac disease and started to develop protocols. However, the difference between the EDTA chelation agents was, and still is, not clear to many chelation therapists.

Since 2000, the chelation therapy movement has gained wide acceptance among environmental physicians and natural healthcare practitioners. In Germany, nonmedical practitioners are fond of using CaEDTA, either as a ‘push’ or a fast infusion, all of which is against protocol. Few are aware of the risks involved. The fact that NaMgEDTA should not be administered to children, and that (however few) mortalities happened is overlooked. Pharmacies provide EDTAs without Rx to nonmedical professionals and without proper drug information.

In 2003, the FDA responded to the death of a child when a physician wrongly applied Na$_2$EDTA instead of CaEDTA, a true medical error [3]. CaEDTA has been approved for the treatment of acute lead intoxication, and while the FDA has issued a public health advisory around reported accidents following the administration of ‘EDTA’ it clearly states that “Seven of the 11 deaths resulted from confusion of edetate disodium with another drug. In five cases, edetate disodium was administered instead of edetate calcium disodium. In two cases, edetate disodium was administered instead of the drug. Etoimidate. Etoimidate is not a form of EDTA”[3].

With this information, we aim to clarify the different mode of action of the EDTAs and their appropriate medical use. In Germany, nonmedical practitioners widely use EDTA chelation, not being aware of the differences of the various substances. In the USA, medical practitioners promote EDTA chelation, often as an alternative to conventional treatments for a variety of chronic diseases and these easily accessed website information are used as teaching material. Some websites promote the administration of the "CaEDTA push", which is clearly against protocol, increasing the risk of iatrogenic accidents.
Materials and Method

To analytically demonstrate the differences and similarities of the chelating agents Na₂EDTA and CaEDTA we statistically evaluated our data base. Over the past 10 years, we methodically supplied physicians with detailed information about chelating agents and proper protocols, asked clinics to submit samples with treatment details, including amounts of chelating agents used, and patient history. We provided urine sample collection protocols that all proper information regarding metal binding and urinary excretion.

For this review, the urine data utilized is based on samples received during 2014-2015 from mostly German chelation therapists. Protocol instructions, including sampling instructions were provided. The samples included in our study are from chronically exposed adult patients. Acutely intoxicated patients were not included. To avoid external contamination, samples were collected into metal-free tubes, provided by the laboratory. Samples were shipped to the laboratory via regular post or courier.

Sample Testing

For urine sample digestion the following method was used:

- 500 µL Urine were pipetted in a 15 ml tube
- +50 µL of Internal Standard Solution was added (Sc, Y, Ho) à 200 patients. Acutely intoxicated patients were not included. To avoid external contamination, samples were collected into metal-free tubes, provided by the laboratory. Samples were shipped to the laboratory via regular post or courier.

Urine metal analysis was performed using the 7700 Series Inductively Coupled Mass Spectrophotometer (ICP-MS) with Agilent’s Octopole Reaction System (ORS), an improved type of mass spectrometer, which provides sensitive, robust, interference-free analysis of difficult, high-matrix samples. With five times the sensitivity of its predecessor and increased matrix tolerance, the ORS system replaces both GFAA and ICP-OES instruments in addition to older generation ICP-MS systems [4].

![Image of a table]

<table>
<thead>
<tr>
<th># of tests</th>
<th>Cd</th>
<th>Cu</th>
<th>Fe</th>
<th>Ni</th>
<th>Lead</th>
<th>Mn</th>
<th>Mo</th>
<th>Zn</th>
</tr>
</thead>
<tbody>
<tr>
<td>2618 Baseline urine</td>
<td>0.29</td>
<td>8.95</td>
<td>23.56</td>
<td>6.04</td>
<td>1.31</td>
<td>2.94</td>
<td>2.94</td>
<td>0.44</td>
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<tr>
<td>93 CaEDTA 1.9 g</td>
<td>1.03</td>
<td>24.59</td>
<td>283.92</td>
<td>9.52</td>
<td>10.25</td>
<td>40.13</td>
<td>24.26</td>
<td>14.16</td>
</tr>
<tr>
<td>131 NaMgEDTA 3 g</td>
<td>1.06</td>
<td>62.74</td>
<td>282.26</td>
<td>9.81</td>
<td>17.85</td>
<td>27.93</td>
<td>23.29</td>
<td>14.04</td>
</tr>
</tbody>
</table>

Source: MTM laboratory data base 2015

Table 1: Na₂EDTA and NaCaEDTA Comparison of Urine Excretion mean values after chelation.

![Image of a table]

<table>
<thead>
<tr>
<th># of tests</th>
<th>As</th>
<th>Ba</th>
<th>Hg</th>
<th>Pd</th>
<th>Sb</th>
<th>Se</th>
</tr>
</thead>
<tbody>
<tr>
<td>2618 Baseline urine</td>
<td>26.19</td>
<td>5.89</td>
<td>0.44</td>
<td>0.27</td>
<td>0.09</td>
<td>22</td>
</tr>
<tr>
<td>93 CaEDTA 1.9 g</td>
<td>21.43</td>
<td>5.02</td>
<td>0.87</td>
<td>0.27</td>
<td>0.1</td>
<td>24.2</td>
</tr>
<tr>
<td>131 NaMgEDTA 3 g</td>
<td>16.02</td>
<td>4.23</td>
<td>1.17</td>
<td>0.2</td>
<td>0.19</td>
<td>27.32</td>
</tr>
</tbody>
</table>

Source: MTM laboratory data base 2015

Table 2: Na₂EDTA and NaCaEDTA Comparison of Urine Excretion mean values after chelation.
NaCiccone MM et al. suggest that treatment with Beniparin is a safe and effective alternative to conventional anticoagulants, particularly for deep vein thrombosis. When it comes to the binding of metals, CaEDTA, Na\textsubscript{2}EDTA, NaMgEDTA was used compared to 1.9 g of CaEDTA. However, the mean value for mercury increased, but not to the level of a DMPS (2,3-Dimercapto-1-propanesulfonic acid) provocation test. Our statistical evaluation of 2659 intravenously administered DMPS tests (1 Ampule Dimaval = 250 mg DMPS) showed a mean mercury value of 16.7 mcg/g creatinine, compared to the EDTA mean mercury value of around 1 mcg/g creatinine (Table 2). Again, none of the data involved came from acutely intoxicated patients.

### Differences of CaEDTA and Na\textsubscript{2}EDTA

As pointed out before, CaEDTA is bound to Calcium, while Na\textsubscript{2}EDTA or NaMgEDTA is not. One 5 ml ampule CaEDTA, containing 1.9 g CaEDTA releases about 203 mg Calcium into the blood stream. The Textbook of Pediatric Medicine warns that "Side effects of Na\textsubscript{2}EDTA include local reactions at injection sites, fever, hypercalcemia" [8].

Na\textsubscript{2}EDTA or NaMgEDTA only bind calcium. Figure 1 indicates that the intravenous administration of 1.9 g CaEDTA, either as a 'push' or short infusion) increased the urine calcium concentration above 400 mg/g creatinine. The infusion of 3 g NaMgEDTA in a 500 ml carrier solution released no calcium, but resulted in calcium binding. The urine calcium concentration following the administration of 3 g NaMgEDTA is nearly the same as that from the administration of 1.9 g CaEDTA.

Patients suffering from hyperparathyroidism should not be treated with CaEDTA. Primary hyperparathyroidism occurs in 25 per 100,000 persons in the general population and in 75 per 100,000 hospitalized patients.

Hypercalcemia is a fairly common metabolic emergency. Between 20% and 40% of patients with cancer develop hypercalcemia at some point in their disease (this may be decreasing with the use of bisphosphates, but data are lacking). Because hypercalcemia is associated with malignancies and not uncommonly found in cancer patients, CaEDTA chelation should not be an option for the treatment of cancer [9].

### Na\textsubscript{2}EDTA and Hypocalcemia

Because of its strong calcium binding ability, Na\textsubscript{2}EDTA was approved to treat digitalis intoxication. In the 1950s, the FDA also approved Na\textsubscript{2}EDTA for the emergency lowering of hypercalcemia, a rare disease.

Na\textsubscript{2}EDTA infusions cause a temporary lowering of serum calcium levels. Patients with parathyroid disorders and particularly children are at risk of developing hypocalcemia during treatment. Consequently, Na\textsubscript{2}EDTA or NaMgEDTA infusions must be administered slowly.

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**Table 3: Na\textsubscript{2}EDTA and NaCaEDTA Comparison of Urine Excretion mean values after chelation.**

<table>
<thead>
<tr>
<th># of tests</th>
<th>Sn</th>
<th>Sr</th>
<th>Ti</th>
<th>U</th>
<th>W</th>
</tr>
</thead>
<tbody>
<tr>
<td>2618</td>
<td>0.57</td>
<td>150</td>
<td>0.25</td>
<td>0.07</td>
<td>0.17</td>
</tr>
<tr>
<td>93</td>
<td>0.44</td>
<td>147</td>
<td>0.22</td>
<td>0.05</td>
<td>0.13</td>
</tr>
<tr>
<td>131</td>
<td>0.59</td>
<td>152</td>
<td>0.27</td>
<td>0.01</td>
<td>0.11</td>
</tr>
</tbody>
</table>

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**Figure 1:** Comparison of mean urinary calcium value after Na\textsubscript{2}EDTA and CaEDTA after chelation.

NaMgEDTA is also effective in improving vasodilation, however Ciccone MM et al. suggest that treatment with Beniparin is a safe and effective alternative treatment, especially for deep vein thrombosis. CaEDTA does not noticeably improve vasodilation, and it does not bind calcium. When it comes to binding lead or other potentially toxic metals, CaEDTA, Na\textsubscript{2}EDTA and NaMgEDTA show similar metal binding abilities. Na\textsubscript{2}EDTA is rarely administered without the addition of magnesium, hence our data bank not have sufficient data to include Na\textsubscript{2}EDTA in the comparison as outlined below (Figure 1, Tables 1-3).

Table 1 indicates that CaEDTA and NaMgEDTA are useful for the chelation of cadmium (Cd). Urine cadmium concentration after provocation with NaMgEDTA and CaEDTA show equal results for cadmium.

The binding of nickel (Ni) may be considered marginal for both chelating agents. For the chelation of lead (Pb), CaEDTA is generally preferred, however our data indicates that NaMgEDTA may also be considered for the detoxification of lead-intoxicated adults. (Caution: because of the strong metal binding, Na\textsubscript{2}EDTA and NaMgEDTA should not be used for the treatment of children!).

The highlighted mean values for the elements copper (Cu), Lead (Pb) and manganese (Mn) may be due to the fact that 3 g of NaMgEDTA was used compared to 1.9 g of CaEDTA. The 5 ml Na\textsubscript{2}EDTA Ampules generally supply 3 g of Na\textsubscript{2}EDTA while the 5 ml CaEDTA supplied by pharmacies contain 1.9 g CaEDTA.

It should be of interest to physician involved in environmental medicine that urine provocation with CaEDTA or NaMgEDTA did not result in an increase in metal binding for the elements As (Arsenic), Ba (Barium), Pd (Palladium), Sb (Antimon), Se (Selenium), Sn (Tin), Sr (Strontium), Tl (Thallium), U (Uranium) and W (Tungsten).

It should be noted that both CaEDTA and NaMgEDTA are not approved to treat digitalis intoxication. In the 1950s, the FDA also approved Na\textsubscript{2}EDTA for the emergency lowering of hypercalcemia, a rare disease.

Na\textsubscript{2}EDTA infusions cause a temporary lowering of serum calcium levels. Patients with parathyroid disorders and particularly children are at risk of developing hypocalcemia during treatment. Consequently, Na\textsubscript{2}EDTA or NaMgEDTA infusions must be administered slowly.
Case Report

Texas: In February 2005, a girl aged 2 years who was tested for blood lead during routine health surveillance had a capillary BLL (blood lead level) of 47 µg/dL. A venous BLL of 48 µg/dL obtained 12 days later confirmed the elevated BLL. A complete blood count and iron study conducted concurrently revealed low serum iron levels and borderline anemia. On February 28, 2005, the girl was admitted to a local medical center for combined oral and IV chelation therapy.

The patient’s blood electrolytes at admission were within normal limits. Initial medication orders included IV Na₂EDTA and oral Sucrimer (DMSA (Meso-2-3-Dimercaptosuccinic acid), the agent primarily used for treatment of lead poisoning in children). The medication order was corrected by the pediatric resident to IV CaEDTA.

At 4:00 p.m. on the day of admission, the patient received her first dose of IV CaEDTA (300 mg in 100 ml normal saline at 25 ml/h). At 4:35 p.m., she was administered 200 mg of oral Sucrimer. Her vital signs remained normal throughout the night. At 4:00 a.m. the next day, a dose of IV Na₂EDTA (instead of IV CaEDTA) was administered. An hour later, the patient’s serum calcium had decreased to 5.2 mg/dL (normal value for pediatric patients: 8.5-10.5 mg/dL). At 7:05 a.m., the child’s mother noticed that the child was limp and not breathing. Bedside procedures did not restore a normal cardiac rhythm, and a cardiac resuscitation code was called at 7:25 a.m. Blood levels were obtained at 8:12 a.m. from the resuscitation failed, and the girl was pronounced dead.

The cause of death was recorded as sudden cardiac arrest resulting from hypocalcemia associated with chelation therapy. The hospital's child mortality review board findings indicated that a dose of IV Na₂EDTA was unintentionally administered to the child [9].

NaMgEDTA, Diabetes and Vascular Disease

It is estimated that during the past 50 years, over one million patients have received intravenous chelation therapy with NaMgEDTA. Atherosclerotic patients improved, which surprised even those who had administered the infusions. Consequently, the American College of Advanced Medicine introduced thousands of medical doctors to NaMgEDTA chelation therapy. courses were taught, attracting physicians from around the world. NaMgEDTA infusions are now used around the globe, largely for the treatment of cardiovascular disease or vascular problems as seen in diabetics.

The role of NaMgEDTA Chelation therapy in Diabetes and Heart Disease has long been debated. A Google search leads to nearly 500,000 pages of pro and con information. To prove or disprove NaMgEDTAs use in cardiovascular medicine, the National Institutes of Health has spent $30 million on a major clinical trial. The study was finalized in 2013. The chief investigator Prof. Gervasio A Lamas MD from Mount Sinai Medical Center found a significant 15% decrease in risk of the primary composite end point among diabetes patients. NaMgEDTA treatment caused a 40% reduction in total mortality, a 40% reduction in recurrent MI, and about a 50% reduction in mortality in patients with diabetes [10].

There are several mechanisms known that benefit cardiovascular disease patients. Research published by Yasuyuki Fujiwara of the Department of Environmental Health, Faculty of Pharmaceutical Sciences, Hokuriku University, Japan indicates that lead and cadmium inhibit repair of vascular cells. EDTA (CaEDTA, Na₂EDTA, NaMgEDTA) effectively removes lead and cadmium [11].

Summary and Conclusion

Both of the chelating agents, CaEDTA and NaMgEDTA, can be used for the detoxification of Cadmium, Copper, Iron, Nickel, Lead and Zinc. Because of their chemical and pharmacological difference, the agents differently affect the calcium metabolism. NaMgEDTA or NaMgEDTA infusions cause a temporary lowering of serum calcium levels, which can be life-threatening if used in children.

From our data we suggest that neither CaEDTA nor NaMgEDTA are useful for the detoxification of As (Arsenic), Ba (Barium), Pd (Palladium), Sb (Antimon), Se (Selenium), Sn (Tin), Sr (Strontium), Tl (Thallium), U (Uranium) and W (Tungsten). The EDTAs Mercury binding ability is not promising.

All the EDTAs should be used with care. One consequence of administering CaEDTA is the release of calcium into the bloodstream, which may lead to hypercalcemia, a condition associated with malignancies.

In Germany, the EDTAs are listed as prescription items (Rx) and thus are officially available to medical physicians only. However, off-label, pharmacies provide these Rx drugs to nonmedical practitioners who generally have little or no training in parenteral applications of such chemicals and are largely unaware of the pharmacological differences of the EDTA chelating agents discussed here. We recommend that teaching institutions provide more thorough information regarding the use of these chelating agents. We also recommend that pharmacists and internet pharmacies are more closely regulated when supplying the EDTAs.

References

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