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High-Strength Low-Alloy (HSLA) Mg–Zn–Ca Alloys with Excellent Biodegradation Performance

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This article deals with the development of fine-grained high-strength low-alloy (HSLA) magnesium alloys intended for use as biodegradable implant material. The alloys contain solely low amounts of Zn and Ca as alloying elements. We illustrate the development path starting from the high-Zn-containing ZX50 (MgZn5Ca0.25) alloy with conventional purity, to an ultrahigh-purity ZX50 modification, and further to the ultrahigh-purity Zn-lean alloy ZX10 (MgZn1Ca0.3). It is shown that alloys with high Zn-content are prone to biocorrosion in various environments, most probably because of the presence of the intermetallic phase $Mg_6Zn_3Ca_2$. A reduction of the Zn content results in (Mg,Zn)₂Ca phase formation. This phase is less noble than the Mg-matrix and therefore, in contrast to Mg₆Zn₃Ca₂, does not act as cathodic site. A finegrained microstructure is achieved by the controlled formation of fine and homogeneously distributed (Mg,Zn)₂Ca precipitates, which influence dynamic recrystallization and grain growth during hot forming. Such design scheme is comparable to that of HSLA steels, where low amounts of alloying elements are intended to produce a very fine dispersion of particles to increase the material's strength by refining the grain size. Consequently our new, ultrapure ZX10 alloy exhibits high strength (yield strength $\dot{R}_{\rm p}$ = 240 MPa, ultimate tensile strength $R_{\rm m}$ = 255 MPa) and simultaneously high ductility (elongation to fracture A = 27%), as well as low mechanical anisotropy. Because of the anodic nature of the (Mg,Zn)₂Ca particles used in the HSLA concept, the in vivo degradation in a rat femur implantation study is very slow and homogeneous without clinically observable hydrogen evolution, making the ZX10 alloy a promising material for biodegradable implants.

INTRODUCTION

Magnesium (Mg) and its alloys are biocompatible and biodegradable, and thus, these materials are promising for medical applications.^{1–3} For use in such applications, they need to fulfill challenging mechanical, electrochemical, and biological requirements. A recent study has demonstrated that the alloying system Mg–Zn–Ca may offer a very attractive combination of high strength and simultaneously high ductility; $R_{\rm p} > 200$ MPa, $R_{\rm m} > 250$ MPa, $A > 20\%.^4$ From a biochemistry viewpoint, Zn and Ca are both essential trace elements in the human body and, as alloying element in Mg, known to positively influence bone healing and cell reactions.^{5,6}

The highly favorable mechanical properties of these types of Mg alloys can primarily be ascribed to their very fine grain size. High strength is achieved by a high degree of grain-boundary strengthening and by solid-solution hardening of Zn.^{4,7,8} The ductility of fine-grained Mg–Zn–Ca alloys is increased

by the activation of nonbasal slip induced by compatibility stresses at the grain boundaries.^{9,10} Furthermore, a grain size of less than 5 μ m does not only improve strength and ductility but also reduces the mechanical anisotropy.^{4,7,8,11,12}

The most important prerequisite for achieving ultrafine-grained structures in Mg–Zn–Ca alloys is the presence of intermetallic particles (IMPs) of controlled density and morphology. Because both alloying elements participate in the formation of the grain-refining IMPs and Zn also contributes to solid-solution strengthening, an ideal Ca and Zn amount was worked out by Hänzi et al.⁴ to ensure the best possible grain-boundary strengthening and solid-solution hardening. The alloy ZX50 with the composition window 0.2–0.4 wt.% Ca and 5–6 wt.% Zn was found to optimally satisfy the requirements for high strength and simultaneously high ductility.

ALLOY ZX50—FROM "HIGH PURITY" TO "ULTRAHIGH PURITY"

Because of its attractive mechanical properties and absence of rare-earth (RE) elements, a ZX50 alloy with nominally 5 wt.% Zn and 0.25 wt.% Ca and conventional purity (CP) was selected for a long-term animal study and compared to the REcontaining alloy WZ21.¹³ Figure 1 illustrates the degradation performance 4 weeks after implantation. Obviously, the degradation of the CP ZX50 alloy is significantly higher than that of WZ21. From the weight loss data in Ref. 13 the *in vivo* degradation rate P_{iv} can be estimated to be roughly 1.3 mm/year for CP ZX50 and 0.16 mm/year for WZ21. Although the *in vivo* degradation rate of WZ21 is relatively low, the corresponding hydrogen gas formation $(Mg + 2H_2O \rightarrow Mg(OH)_2 + H_2)$ is still too high to be absorbed by the surrounding tissue. This reveals the particular need for further improvement of the degradation performance, but without using RE additions, *nota bene*. For more details on the biological concerns regarding RE elements, see Ref. 14.

Kraus et al.¹³ did not discuss the metal-physical reasons for the high degradation rate of CP ZX50. However, in the current study, we will illustrate that at least two metallurgical aspects are of particular importance: (i) trace elements, in particular iron, exceeding the impurity tolerance limits, which results in the formation of noble Fe-containing intermetallic particles,¹⁵ and (ii) formation of noble Zn-containing intermetallic particles of type $Mg_6Zn_3Ca_2$.¹⁶ Both Fe-containing IMPs and Zncontaining IMPs may act as cathodic sites and lead to anodic dissolution of the Mg matrix.

It is a well-known fact that the impurity elements Fe, Cu, Ni and Co enhance the saltwater corrosion of Mg alloys.¹⁷ The corrosion rates are significantly accelerated when their concentrations exceed certain values.^{15,18} They are particularly harmful because of their low solid solubility and thus can form precipitates, which provide effective cathodic sites even at low concentrations.^{19,20} To check the influence of trace elements on the degradation performance, we prepared ZX50 alloys using high-purity (HP) Mg and vacuum-distilled ultrahigh purity (XHP) Mg.²¹ The chemical composition with specific consideration of "harmful" elements is given in



Fig. 1. µCT reconstructions (two-dimensional slices) of Mg-pins implanted into the femur of Sprague–Dawley rats 4 weeks after operation. The degradation rate of the alloy CP ZX50 (a) is more than 3 times higher than that of WZ21 (b).¹³ Massive hydrogen gas formation affects bone and tissue healing for CP ZX50; only moderate hydrogen pocket formation is visible for WZ21.

Table I. Chemical composition of ZX50 with conventional purity (CP), high-purity (HP), and ultra-high purity (XHP)

Alloy	Zn (wt.%)	Ca (wt.%)	Mn (ppm)	Si (ppm)	Fe (ppm)	Cu (ppm)	Ni (ppm)	Co (ppm)
CP ZX50	5.24	0.27	1630	440	42	9	8	8
HP ZX50	5.21	0.29	160	430	31	8	7	5
XHP ZX50	5.26	0.31	0.8	2.9	0.5	0.09	0.05	< 0.05



Fig. 2. Isopleth plot of Mg-Zn5-Ca0.25-Fe <0-0.01>. The traces of the alloy's constitution points are indicated by the dashed lines; green (XHP ZX50), red (HP ZX50), and magenta (CP ZX50).

Table I. The use of XHP-Mg and ultrapure Zn and Ca (see the "Methods" section) for the XHP ZX50 synthesis results in a reduction of the trace element content by a factor of about 100.

Figure 2 shows an isopleth plot of the Mg-Zn5-Ca0.25-Fe system calculated using Pandat (CompuTherm, Madison, WI), a simulation software for multicomponent phase diagram calculations.²² It can be noted that in solid ZX50, the solubility limit for Fe is extremely low, i.e., less than 1 ppm Fe. Dashed vertical lines indicate the trace of the constitution points of the three alloys: CP ZX50, HP ZX50, and XHP ZX50. Whereas XHP ZX50 is close to the tolerance limit, the alloys CP ZX50 and HP ZX50 are significantly above the limit, and formation of Fe-containing particles (body-centered cubic [BCC]) will take place. In this respect, it is important to note that, according to experimental findings and thermodynamic simulations,²² the BCC phase contains besides Fe mainly Mn and Si, and thus, their volume content is dependent not only on the Fe level but also on the Mn and Si content (see Table I). Because the alloys were processed (i.e., extruded to rods) at temperatures of around 300°C (see the "Methods" section), we calculated the equilibrium mole fraction of BCC at this temperature. With 3.8×10^{-4} BCC in CP ZX50, it is more than one order of magnitude higher than in HP ZX50 (1.6×10^{-5}) and two orders of magnitude higher than in XHP ZX50 ($< 1 \times 10^{-6}$).

All three alloys were tested on their corrosion/ degradation behavior in simulated body fluid (SBF). Either carbon dioxide, CO₂, or Tris, i.e., tris-(hydroxymethyl)aminomethane, (HOCH₂)₃CNH₂, was used as buffer. The degradation was measured by the hydrogen evolution method²³ and evaluated using an improved testing setup.²⁴ Figure 3 depicts the mean hydrogen evolution in CO₂-buffered SBF versus time of the ZX50 alloys. The amounts of hydrogen increase steadily at the beginning with a decrease in degradation rate with increasing time.



Fig. 3. Mean hydrogen evolution versus time of CP, HP, and XHP ZX50 immersed in CO_2 -buffered SBF with pH 7.45 at 37°C. (Note the different scales at the *y*-axis; typical data scatter is illustrated by error bars).

As expected, the XHP ZX50 alloy evolves significantly less hydrogen than the CP and HP alloys within the same time.

This significant improvement (after 2 weeks by more than an order of magnitude from CP to HP, and a factor of ≈ 3 from HP to XHP) can be attributed to the reduced trace element content below or close to the impurity tolerance limit and the related avoidance of degradation-generating cathodic sites. Very recently, a similar effect of purity was observed by Cao et al.²⁵ using unalloyed XHP Mg²¹ in 3.5% NaCl solution. Hence, the XHP strategy can be assumed to be effective for various Mg-based alloying systems. However, as shown in the following, important limitations have to be considered regarding the corrosive environment.

First, we evaluate the influence of the buffering media. Figure 4 shows the ratio of released hydrogen in Tris- and CO_2 -buffered SBF (H₂(Tris)/ $H_2(CO_2)$) at 250 h of immersion for HP and XHP Mg, and the Y-containing alloys WE43²⁶ and WZ21.²⁷ Data for the modified alloy XHP ZX10 (MgZn1Ca0.3) are also included; the design strategy of this alloy will be discussed in detail in the next section. Changing from CO₂ buffer to the organic compound Tris results in an enormous increase in the degradation rate. Such effect of organic buffering agents is well known for various Mg alloys,^{24,28,29} but for ZX50, in both conditions HP and XHP, it is extraordinarily pronounced (note that CP ZX50 behaves similar but is not further considered because of its high degradation rate already in CO_2 -buffered SBF). Obviously, the high-Zn-containing alloys react very sensitively to the presence of the organic buffer agent. The degradation of HP and XHP ZX50 in Tris is extremely fast and could not be measured properly in our test device. Compared to ZX50, the Y-containing alloys WE43 and ZW21 show a significantly lower but still



Fig. 4. Ratio of released hydrogen in Tris- and CO₂-buffered SBF at 250 h of immersion. Pure Mg shows only moderate dependency on the buffer. Stronger dependence is observed for the RE-containing alloys WE43 and WZ21. For HP ZX50 and XHP ZX50, the degradation in Tris-buffered SBF is extremely high and too fast to be measured properly. A reduction of the Zn content reduces the dependency on the buffer very strongly, as demonstrated for the alloy ZX10.

considerable acceleration of the degradation, whereas pure Mg exhibits only moderate dependency on the buffer.

The strong acceleration in Tris-buffered SBF is accompanied by a change in the character of the corrosion morphology (Fig. 5). For the alloys in CO₂buffered SBF, a very homogeneous degradation can be noticed, whereas in Tris-SBF, a distinct localized corrosion attack is observed, and the local attack is most pronounced for the fast-degrading alloys ZX50. However, such change in the degradation morphology does not take place in unalloyed Mg. Furthermore, only very weak localized attack is notable on the alloy XHP ZX10. These findings suggest a strong influence of the alloy's chemical composition on the degradation susceptibility in specific environments.

Recently, several studies suggested that SBF with CO_2 buffer simulates the *in vivo* situation better than organic buffer agents like Tris.^{24,30,31} However, it has become apparent that this is not the case for the high-Zn containing ZX50 alloys. Unpublished immersion tests in protein-containing environments and a 3-month animal test performed by our industrial partner have demonstrated that this alloying system degrades significantly faster than when immersed in CO_2 -buffered SBF. Clearly, this result puts into question whether CO_2 really mimics the *in vivo* situation, as it was stated in Refs. 24, 29, and 32 but also it encouraged us to modify our design concept toward Mg–Zn–Ca alloys with lower Zn content, as discussed in the next section.

ALLOY ZX10—CHOICE OF Zn CONTENT

Recently, several studies considered the influence of the Zn content in Mg–Zn–(Ca) alloys on the corrosion behavior in various environments.^{16,33-38} They all emphasize that well-balanced Zn and Ca contents are required to avoid the formation of noble microcathodic IMPs. In particular, Bakhsheshi-Rad et al.^{16,35} pointed out that a too high Zn content is detrimental because of the formation of high amounts of Mg₆Zn₃Ca₂ and Mg₅₁Ca₂₀. Consequently, reduction of the Zn content might be an effective recipe for achieving better biocorrosion resistance. Such a simple procedure, however, would result in significant loss in strength because of reduced solid-solution hardening. From the data in Ref. 4, the solid-solution hardening of Zn can easily be calculated to be roughly 20 MPa/wt.%, and thus, a reduction of the Zn-content from 5% to 1% would result in an undesired loss in yield strength of about 80 MPa. Hence, further measures have to be taken to compensate the reduced solid-solution hardening. According to Refs. 39 and 40, the Hall-Petch coefficient for fine-grained Mg alloys is about 5 MPa mm $^{1/2}$. Accordingly, a reduction in grain size from 4 μ m (grain size of CP ZX50 used in Ref. 13) to 1 μ m would lead to an increase in yield strength of roughly 80 MPa. From these simple calculations, it becomes evident that fine-grained microstructures $(D < 5 \ \mu m)$ are required to keep the yield strength at the desired high level, i.e., above 200 MPa.

The strategy for achieving very fine grains is based on the assumption that intermetallic particles present in the alloy during hot forming, in our case extrusion, might act as obstacles against grain growth according to Zener.⁴¹ However, with respect to the biocorrosion performance we need to ensure that such IMPs do not act as cathodic sites. Südholz et al.⁴² investigated relevant IMPs and found all to be cathodic to Mg, with the exception of Mg₂Ca. This IMP exhibits some solubility for Zn what further allows us to adjust its electrochemical potential. As demonstrated by Cha et al.,³⁴ the addition of Zn to Mg₂Ca shifts the open circuit potential of $(Mg,Zn)_2$ Ca up to that of the Mg matrix, which prevents selective corrosion and induces uniform corrosion.

Figure 6a illustrates the constitution of Mg–Zn– Ca alloys at 300°C, i.e., the temperature chosen for the hot-forming procedure (see the "Methods" section). Because the intention is to avoid the formation of the ternary Mg₆Zn₃Ca₂ phase, the Zn content must be kept below ~ 1.0 wt.%. To benefit from some solid-solution hardening, we chose 1 wt.% Zn and further evaluated the optimal Ca content. First, we considered the grain-refining effect of Mg₂Ca particles. Their Zener pinning pressure increases with the volume fraction and the reduction of particle size, restricting the matrix grain size to a critical maximum (Zener limit).43 Because the volume fraction increases proportionally with the Ca content, a high Ca content would be beneficial. However, to create a very fine particle size, we need to apply a solution treatment prior to the extrusion process, followed by an ageing procedure below or at



Fig. 5. Surface topography of samples after immersion in SBF. (a) XHP ZX50 shows no local corrosion attack after 2 weeks of immersion in CO₂-buffered SBF and (b) massive localized attack in Tris-buffered SBF already after 3 days. (c) No indication of local corrosion attack is visible for XHP ZX10 after 2 weeks of immersion in Tris-buffered SBF.

the hot-forming temperature (see the "Methods" section). The insert in Fig. 6b shows the width of the solution-treatment window ΔT_{ST} (Zn and Ca are completely dissolved in Mg and the solidus temperature is still not reached, see also Fig. 6b) at a Zn content of 1 wt.%. Obviously, with respect to process reliability, the Ca content must be kept significantly below 0.4 wt.%. A compromise of 0.3 wt.% Ca has proven to be favorable. The constitution of the Znlean alloy MgZn1Ca0.3 (ZX10) is shown in Fig. 6b. As required, only the Mg₂Ca phase is present at the processing temperature of 300°C. According to its Zener drag effect, the resulting grain size stays at low levels, i.e., $D \approx 2 \ \mu m$ (for extrusion parameters, see the "Methods" section). The corresponding mechanical properties of this newly developed alloy, exhibiting the lowest content of alloving elements of all technical Mg alloys, are as follows. In tension: yield strength TYS = 240 MPa, ultimate tensile strength UTS = 255 MPa, and elongation to fracture A = 27%; in compression: CYS = 205 MPa, UCS = 245 MPa, and A = 13%. These values show that the alloy features not only high strength and good ductility but also low mechanical anisotropy.

Interestingly, our design strategy is comparable to HSLA steels, where low amounts of alloying elements are intended to produce a very fine dispersion of carbides, which increase the material's strength by refining the grain size as a result of the effect of particles on dynamic recrystallization and grain growth.⁴⁴

The good mechanical properties of the alloy XHP ZX10 are accompanied by excellent biocorrosion behavior. Figure 7 illustrates its *in vivo* degradation performance compared to CP ZX50 after 12 weeks of implantation time (rat femur implantation of pins with 1.6 mm diameter). The fast corroding CP ZX50 has completely degraded,¹³ while the newly developed XHP ZX10 exhibits the desired slow and

homogeneous degradation. It is important to note that no hydrogen bubbles can be observed. Apparently, the living organism was able to absorb the generated low amount of hydrogen; this fact is of high relevance from a medical point of view. Details of the progress of biodegradation and the corresponding bone and tissue response will be reported elsewhere.

SUMMARY AND CONCLUSIONS

We have reported on the development of a biocompatible, HSLA magnesium alloy for use as biodegradable implant material. Following the pathway from a high-Zn containing alloy (ZX50) and reducing the impurities of corrosion-provoking elements (such as Fe, Cu, Ni, Co, Mn, and Si) down to the <3 ppm level, we developed a new XHP Mg alloy (ZX10, MgZn1Ca0.3 in wt.% or Mg_{99.45}Zn_{0.35}Ca_{0.2} in at.%), which contains only 0.55 at.% of alloying elements. The reduction of the Zn content generates after processing the formation of a (Mg,Zn)₂Ca phase, which can be homogeneously distributed as nanometer-sized IMPs in the Mg matrix.

The new Mg alloy ZX10 shows a very fine microstructure for a technical wrought alloy, with a matrix grain size of $\approx 2 \mu m$. This small grain size generates outstanding mechanical properties, such as high strength and simultaneously high ductility, as well as low mechanical anisotropy. The (Mg,Zn)₂Ca IMPs are less noble than the matrix and thus can act as anodic sites. As a consequence, the alloy degrades homogeneously in *in vivo* studies (rat femur implantation) and at the desired low level without any hydrogen bubble formation. The superior *in vivo* degradation performance and mechanical properties make the ZX10 alloy the most ideal technical alloy for biodegradable implant applications.







Fig. 7. µCT reconstructions (two-dimensional slices) of Mg pins implanted into the femur of Sprague–Dawley rats 12 weeks after operation. Whereas the 1st-generation alloy CP ZX50 has completely degraded and severe irritation of the bone is observed (a), ¹³ only slight homogeneous degradation is noticed for XHP ZX10 (b). Furthermore, no hydrogen bubbles are observed in the case of the newly developed XHP ZX10 alloy.

METHODS

Preparation of Mg Alloys

For the production of HP ZX50, high-purity Mg (99.98%), Ca (99.0%), and Zn (99.99%) were used, while for the XHP alloys ZX50 and ZX10, distilled ultrapure Mg (99.999%),²¹ Ca (99.99%), and Zn (99.999%) were synthesized in a graphite crucible under a protective gas mixture at 750°C. Subsequently, the melt was poured into a conical graphite mold (average diameter 55 mm, height \approx 150 mm), which was water cooled at the bottom to force directional solidification to avoid shrinkage cavities. Afterward, the billets were homogenized (ZX50 alloys: 350°C/12 h; ZX10 alloy: 350°C/12 h plus 450°C/8 h) followed by cooling with pressurized air. The chemical compositions of the samples were determined by glow discharge mass spectrometry.

Extrusion Procedure

The homogenized alloys were machined to billets, measuring 50 mm in diameter and 120 mm in length. For XHP ZX10, the billet was aged at 250°C for 30 min to create fine and homogeneously distributed intermetallic particles prior to the extrusion process. Afterward, the billet was heated to 300°C and indirect extrusion was performed at a ram speed of 0.15 mm/s to a rod profile with a diameter of 10 mm, corresponding to an extrusion ratio of 25:1. Cooling with pressurized air was applied at the exit side of the extrusion press. For the ZX50 alloys, the extrusion to 10 mm diameter was performed in the direct mode at 325°C.

Immersion Tests

The degradation was measured by the hydrogen evolution method²³ and evaluated using an in-house improved testing setup.²⁴ The device exhibits a high gas volume resolution and was especially designed for slow degrading materials; it accounts also for the daily atmospheric pressure fluctuations and gas dissolution. The specimen sizes were used with surface areas in the range of $259 \pm 1 \text{ mm}^2$ to $626 \pm 6.7 \text{ mm}^2$. Three specimens per batch were tested and their mean hydrogen evolution was evaluated to mL/cm². All data points were corrected by the atmospheric pressure and the gas dissolution. The ion concentration of the SBF used for the

immersion tests is described in Ref. 27; either CO₂ (99.90 vol.%) or Tris was used as buffer. During the degradation test, the pH was set at 7.45 ± 0.017 with a temperature of 36.9 ± 0.30 °C for 2 weeks. The samples were ground on abrasive SiC paper of granularity 4000. Just before immersion, they were polished on a 1 μ m polishing cloth, cleaned in an ultrasonic bath using isopropanol, and dried in hot air.

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