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STRUCTURAL (SUPER)PLASTICITY OF MAGNESIUM MATERIALS

Superplastic behaviour of two magnesium alloys QE22 and EZ33 having a grain size of $\sim 1\mu\text{m}$ was studied at a temperature of 420°C and in the strain rate interval from 1×10^{-4} to $1\times 10^{-3}\text{ s}^{-1}$. The substructure of samples was analysed by the transmission electron microscope before and after deformation. The substructure of samples exhibits an existence of particles in the grain boundaries. These precipitates are very stable even at high deformation temperature and they are reason for the relatively high ductility of alloys. Observed ductility of the WE54/SiCp composite is much lower due to the existence of ceramic particles in the grain boundaries. They inhibit the grain boundary sliding and are the reason for the cavitations and early fracture of the composite.

Keywords: Magnesium alloys, Magnesium alloy based composite, Strain rate sensitivity, Electron microscopy, Superplasticity, Threshold stress

1. Introduction

Low density of magnesium alloys and relatively high specific strength are the main advantages of these materials. Magnesium alloys and composites are used in different industrial applications. As magnesium is recyclable, its extensive usage can minimize the negative impact on the environment due to non-degradable plastic wastes. Magnesium materials have low ductility at ambient temperature. This is a consequence of the hexagonal close-packed (hcp) crystal structure with a limited number of operative slip systems. The plasticity of magnesium alloys and composites increases at elevated temperatures where additional slip systems become active. However, low ductility of magnesium alloys prevents their application hence; the enhanced ductility of magnesium materials is of a special interest [1–4]. It is known that the fine grain size ($d \leq 10\mu\text{m}$) stable at higher temperatures is a main structural requirement for the occurrence of superplasticity. All characteristics of superplasticity (relative elongation to failure ϵ_r , flow stress σ , strain rate sensitivity of flow stress m , optimum strain rate for superplasticity and optimum deformation temperature for superplasticity) depend on the grain size. Fine grain structure with the grain size less than $10\mu\text{m}$ may be obtained by various methods such as phase transformation, recrystallisation, forming of the microduplex structure or severe plastic deformation (equal channel angular pressing, reciprocating extrusion) [5–10]. Structural superplasticity was estimated in several magnesium alloys especially in alloys of the AZ series [11–13] or Mg-Li alloys [14–15]. The superplasticity phenomenon was also observed in ZK60 alloy [16], WE54 alloy [17] and QE22 alloy [18].

In papers [18, 19] the influence of preliminary thermomechanical treatment (hardening, overageing and hot extrusion) on the grain size was investigated. This prior thermomechanical treatment influences the number of recrystallisation nuclei and so the resulting grain size.

The aim of this work is to study the microstructure and substructure of two magnesium alloys containing rare earth elements (RE) and Zr, which exhibited better superplastic properties in comparison with alloys containing Al. The aim is also to find the main features influencing the superplastic behaviour and limiting factors for lower plasticity of particles reinforced composite with the Mg alloy matrix.

2. Materials and experimental methods

QE22 (Mg-2Ag-2RE-Zr), EZ33 (Mg-2.5Zn-3RE-Zr) magnesium alloys and WE54 (Mg-4Y-4RE) reinforced with SiC particles were used in this study. Cylindrical QE22 and EZ33 specimens with a diameter of 72 mm and a length of 80 mm were cut from ingots and exposed to homogenisation treatment at 470°C for 10 h with the subsequent cooling on the air. Due to low diffusion rates of solute atoms in the magnesium matrix the alloys were simultaneously quenched. Precipitation ageing was conducted in the temperature range from 200 to 380°C for 10 hours. After that, the aged specimens were preheated at 350°C and then hot extruded. This device consists of the press with the rated strain of 4000 kN and the tube furnace which heats instrumental rigging

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(container with a diameter of 75 mm, matrix and punch). Hot extrusion was performed in two steps with the extrusion ratio of 20 and 26. Samples for deformation experiments were machined from the extruded bars.

Composite WE54/SiC containing 13 vol. % SiC particles was processed by a powder metallurgy method (PM). Extrusion billets were prepared by mixing of the matrix powder with SiC particles in an asymmetrically moved mixer with subsequent milling in a ball mill. The powder was encapsulated in Mg containers and extruded to rods at 400 °C. The composite samples were not thermally treated. Tensile specimens with a gauge length of 10 mm and a gauge diameter of 6 mm were machined with the tensile axis parallel to the extrusion direction.

The strain rate sensitivity parameter $m = \left(\frac{d \ln \sigma}{d \ln \dot{\epsilon}} \right)_T$ was esti-

mated by the abrupt strain rate changes (SRC) method [20]. SRC tests and tensile tests with a constant strain rate ($\dot{\epsilon}$) were performed at temperatures from 300 to 500 °C and strain rates from 10^{-4} to 10^{-2} s^{-1} . Experiments with the constant strain rate, where the cross-head speed continuously increases, significantly reduces the test time at low strain rates. The best conditions for superplasticity of the alloys were found at a temperature of 420 °C.

Foils for Transmission electron microscopy (TEM) from QE22 and EZ33 alloys were thinned by the electro-polishing method using the TENUPOL 3. The specimens were etched in the solution of 15.9 g of LiCl, 335 g of Mg perchlorate, 1500 ml of methanol, and 300 ml of butoxy-ethanol in the temperature range -55 to -40 °C. After electro-polishing, the foils were washed in methanol. To prevent the formation of an oxide surface layer, TEM foils of all experimental materials had to be stored in vacuum. Investigations of substructure of the materials were carried out with JEOL 2000FX. The transmission electron microscope operating at 200 kV was equipped with an energy dispersive X-ray analyzer (EDAX), which provides the analysis of individual phase composition. Microstructure of WE54/SiC composite was studied using a light microscope Olympus and scanning electron microscope JEOL.

3. Results and discussion

3.1. Microstructure of materials

The grain size of the QE22 and EZ33 alloys was measured in the both as-cast and after thermo-mechanical treatment states. The average grain sizes of the as-cast alloys are higher than 100 μm while in the QE22 and EZ33 alloys after the thermo-mechanical treatment, the grain sizes were reduced to 0.7 μm (QE22) and 1.2 μm (EZ33), respectively.

The substructure of QE22 and EZ33 alloys was investigated in the as-cast and after thermo-mechanical treatment states. Microstructure of the as-cast QE22 alloy consists of α -grains decorated in the grain boundaries by the second phase particles. TEM revealed chains of smaller particles; containing Nd and Ag located at grain

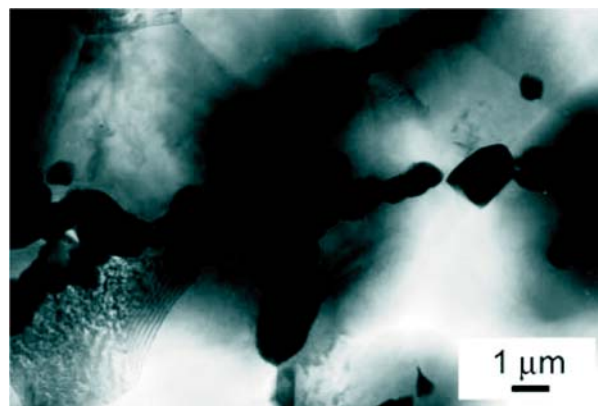


Fig. 1 TEM micrograph of the as-cast QE22 magnesium alloy showing particles of $\text{Mg}_3(\text{Ag}, \text{Nd})$ phase situated in grain boundaries

boundaries (see Fig. 1). These particles were dissolved during homogenisation annealing. TEM showed only groups of smaller and bigger non-dissolved zirconium particles which are visible in Fig. 2a. These Zr particles are very stable and they were not influenced by the homogenisation treatment. Dislocations pinned at

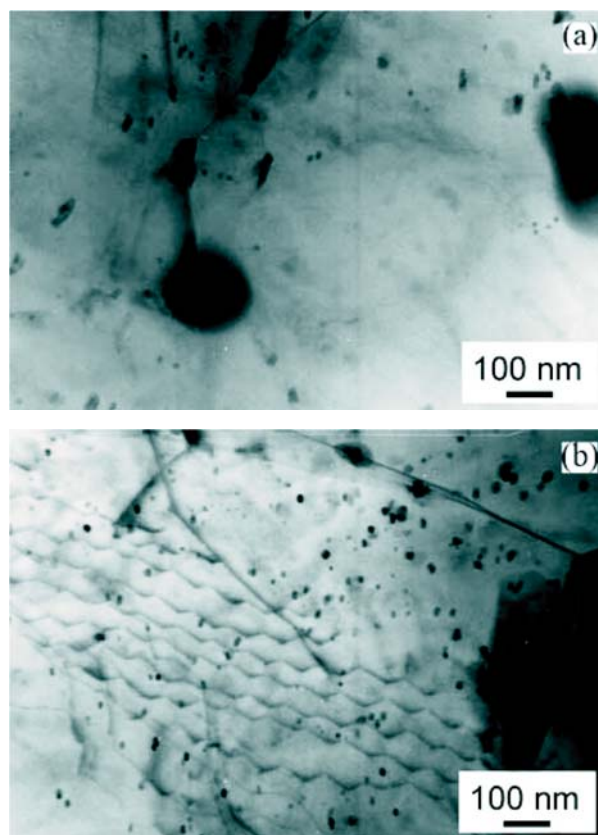


Fig. 2 TEM micrographs of the QE22 magnesium alloy after the homogenisation treatment: (a) fine Zr particles, (b) Zr particles in partially visible dislocation network

Zr particles are also visible in Fig. 2b. After thermo-mechanical treatment (age annealing at 350 °C for 10 h and hot extrusion) the particles at grain boundaries appeared again (see Fig. 3) and they were found again together with the smaller Zr particles also after deformation at 420 °C.

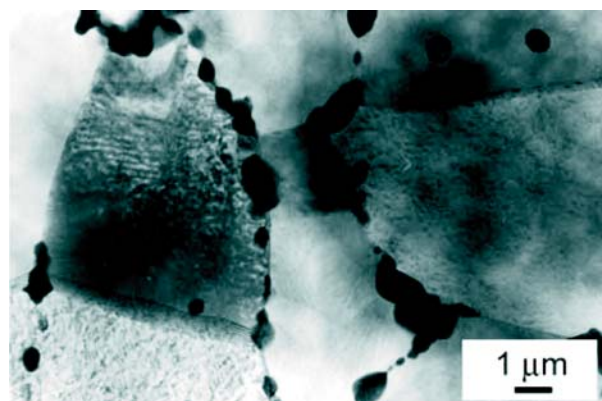


Fig. 3 TEM micrographs of the QE22 magnesium alloy after thermo-mechanical treatment showing chain of particles of $Mg_3(Ag, Nd)$ phase in grain boundaries

The interior of grains after deformation at 420 °C was clear without dislocations. Round particles appearing after the thermo-mechanical treatment were identified as $Mg_3(Ag, Nd)$ phase. There are two kinds of round particles of $Mg_3(Ag, Nd)$ phase: bigger particles with an average diameter of 0.2–0.5 μm lying along the grain boundaries and in the triple points and the smaller particles with an average diameter of 50–100 nm situated in the grains interior. The obtained results of the $Mg_3(Ag, Nd)$ phase distribution agree with results in the paper [21] where similar two particle sizes were found in QE22 alloy after T6 temper (annealing for 5.5 h. at 520 °C and ageing 8 h. at 205 °C). Particles placed at grain boundaries contribute to the microstructure stability hereby they hinder the grain growth during heating in the deformation machine and the self deformation. Precipitates in the grain boundaries were found also after the high temperature deformation practically unaffected.

The electron micrograph of the EZ33 alloy is shown in Figure 4 in the as-cast state. Fine Zr particles are visible. These particles remained unaffected by the thermo-mechanical treatment. The microstructure of EZ33 alloy after thermo-mechanical treatment exhibits small α grains containing fine ZnNd particles in the grain boundaries, very probably particles of T phase (Figure 4a). Wei et al. studied the solidification path and phase constituents of alloys in the magnesium rich corner of the Mg-Zn-misch metal system [22]. They found that the T phase having a c-centred orthorhombic crystal structure exhibits a wide range composition of Mg, Zn and RE depending on the alloy composition. The Zr particles (200–100 nm) were observed in the whole volume. In the sample of EZ33 after high temperature exposition during deformation the same particles were found, i.e. T phase in the grain boundaries and small Zr particles in the grains interior (Fig. 4b). While dis-

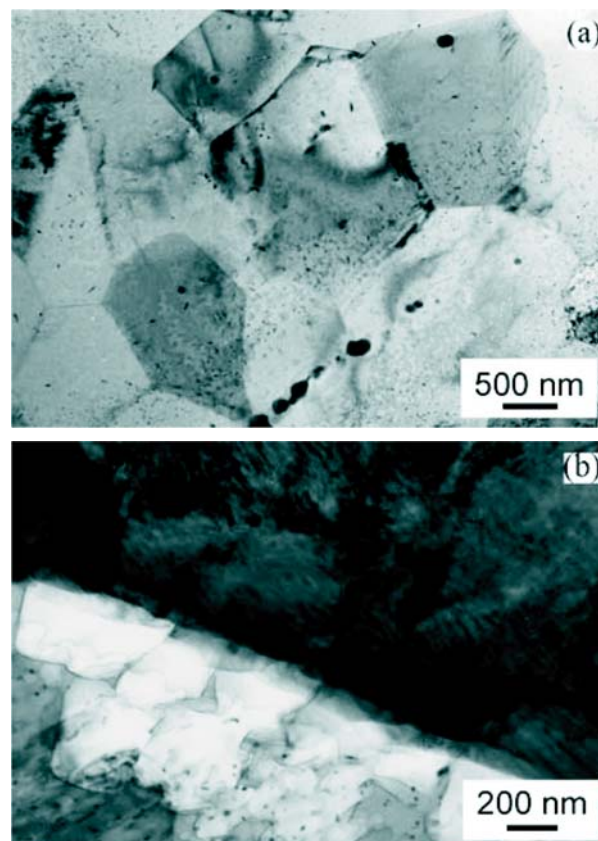


Fig. 4 TEM micrographs of the EZ33 magnesium alloy: after thermomechanical treatment (a) and after high temperature exposition during the deformation process (b)

locations in this material were present after the thermomechanical treatment (see Fig. 5), after the high temperature deformation no dislocations were detected. The average grain size of EZ33 alloy is relatively small (1.2 μm), though it is even smaller (0.7 μm)

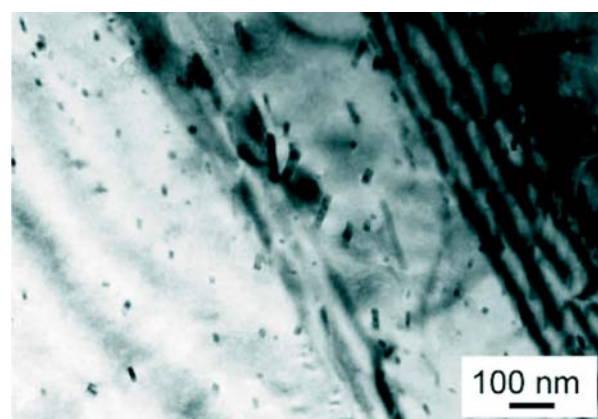


Fig. 5 TEM micrographs of the EZ33 magnesium alloy after thermo-mechanical treatment showing resistance to dislocation motion by the Zr particles

in QE22 alloy. Such a difference between the grain sizes in the alloys can be caused by the presence of alloying element Nd besides Zr. QE22 alloy contains mainly Nd among rare earth elements. Furthermore, fine precipitates of Mg_3 (Ag, Nd) phase are uniformly located also inside of grains. As such Zn (and rare earth) containing alloys are the major Mg alloy group where Zr is used as the grain refiner [23]. Although Zr has long been used as a notable grain refiner for Mg alloys [24] the mechanism of grain refinement by Zr is not yet completely understood. In the alloys, which do not contain Al, small Zr particles remain unsolved in the α -matrix and influence the grain nucleation during solidification or recrystallisation.

Figure 6 shows the microstructure of the WE54/SiC_p composite. The as-extruded bar appears mainly constituted of very small equiaxed grains. The corresponding average grain size was found to be about $\sim 3 \mu m$. The SiC particles (mean particle size $\sim 9 \mu m$) were not uniformly distributed in the matrix; in many cases they formed clusters.

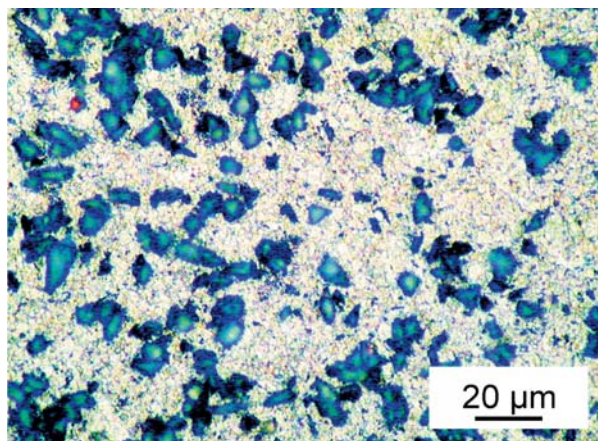


Fig. 6 Microstructure of WE54/SiC_p composite

3.2. Superplastic behaviour of alloys

Values of the elongation to failure ϵ_f introduced in Table 1 together with values of m -parameter were estimated at 420 °C. The highest values of the elongation to fracture $\epsilon_f = 780\%$ (QE22) and 710% (EZ33) were estimated at a strain rate of $\dot{\epsilon} = 3 \times 10^{-4} s^{-1}$. These values correspond to the highest values of the strain rate sensitivity $m = 0.75$ (QE22) and 0.73 (EZ33) which were estimated at the same deformation conditions. As it follows from Table 1, the tensile elongations of all samples exceeded 200%, which represents a substantial improvement over the poor room-temperature ductility, typical for magnesium alloys. The strain rate of materials deformed at high temperatures is done by a relationship of the following form:

$$\dot{\epsilon} = A \frac{Gb}{kT} \left(\frac{b}{d} \right)^n \left(\frac{\sigma}{G} \right)^n D, \quad (1)$$

where A is a dimensionless material constant, G is the shear modulus, b is the Burgers vector, d the grain size, d is the grain size exponent, $n = 1/m$ is the true stress exponent. kT has its usual meaning.

Elongation to failure and strain rate sensitivity values estimated at 420 °C

Table 1

$\dot{\epsilon} (s^{-1})$	QE22		EZ33	
	$\epsilon_f (\%)$	m	$\epsilon_f (\%)$	m
1×10^{-4}	720	0.62	700	0.65
3×10^{-4}	780	0.75	710	0.73
5×10^{-4}	450	0.50	420	0.50
1×10^{-3}	360	0.42	340	0.48
5×10^{-3}	300	0.38	280	0.35
0.01	240	0.32	230	0.30

D is the appropriate diffusion coefficient ($= D_0 \exp(-Q/RT)$, where D_0 the frequency factor, Q the activation energy for the diffusion process and R the gas constant). The steady state plastic flow of coarse-grained metals at high temperatures, above $0.4T_M$ (T_M being the melting temperature), is usually described by dislocation motion and storage on obstacles. These obstacles may be either of non-dislocation type or of dislocation type. Grain boundaries or incoherent precipitates are considered as the obstacles of the non-dislocation type. Dislocation type obstacles are formed by reactions between various types of dislocations. Steady state character of deformation is provided with the dislocation cross slip and also by climbing of dislocation edge segments. In the fine grained materials two other independent mechanisms – grain boundary sliding accommodated by slip and directional diffusional flow may be considered [25-27]. Each of the three deformation mechanisms has specific values of n , p and Q by which the mechanism can be uniquely defined. For the plastic flow by slip of dislocation the high values of the stress exponent ($n = 5$ or higher) are characteristic and the activation energy is close to values for the activation energy of the lattice diffusion or for the pipe diffusion. The grain size exponent p is close to zero [28]. Grain boundary sliding accommodated by dislocation slip or diffusional flow is characterised by the low values of the stress exponent (lower than 3, in the case of ideal superplasticity 1) and the activation energy is equal to the lattice or grain boundary diffusion or combination both. As no dislocations were observed by means of TEM and the estimated values of the stress sensitivity parameter are 1.33 (QE22) and 1.37 (EZ33) we may conclude that the main deformation mechanism during the superplastic flow is the grain boundary sliding accommodated by lattice and grain boundary diffusion.

3.3 Plasticity of WE54/SiC_p composite.

The strain rate sensitivity parameter m was measured in SRC tests. The values of m estimated for different strain rates and tem-

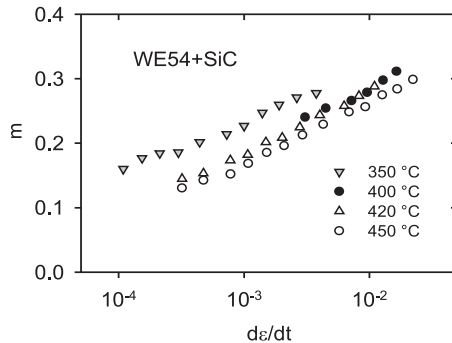


Fig. 7 Strain rate dependence of m -parameter

peratures are introduced in Fig. 7. The strong strain rate dependence of the m -parameter is obvious from the figure. With increasing temperature the dependence is shifted to the higher strain-rates. However, the m parameter slightly increases with temperature; the maximum values are in the vicinity of 0.3 as it is obvious from Table 2. The maximum recorded elongation to failure was 99%, which shows an evidence of the enhanced plasticity, nevertheless this value remains below the bottom limit for superplastic region. The SEM micrograph of the sample exhibiting the highest elongation is shown in Fig. 8 documenting the microstructure after deformation at 450 °C. Numerous cavities formed during the high temperature deformation are visible in Fig. 8. The cavitation primarily occurred at the reinforcement/matrix interfaces which are

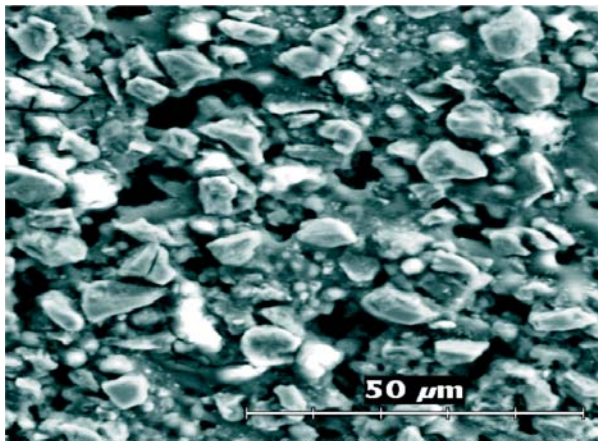


Fig. 8 Micrograph showing cavities formed during high temperature deformation

the preferential sites for the nucleation of cavities. The density of cavities was higher near the fracture surface where elongated cavities were found. Since many of these cavities were fairly large, it is reasonable to assume that the growth and subsequent coalescence and interlinkage of the cavities led to a premature failure.

As it is obvious from Fig. 8 the m -values decrease with decreasing strain rate at low strain rates down to ~ 0.1 . Low m -values in

Table 2
Maximum values of m -parameter and elongation to failure estimated for three temperatures and corresponding strain rates for WE54/SiC composite

Temperature	m_{max}	$\dot{\epsilon}$ (s^{-1})	ϵ_f (%)	σ_{th} (MPa)
350 °C	0.28	4×10^{-3}	66	11.2
420 °C	0.30	2×10^{-2}	95	6.6
450 °C	0.29	5×10^{-2}	99	4.7

the low strain rate regime were generally reported for PM magnesium alloys and Mg based composites [29–32]; they are probably associated with the existence of the threshold stresses σ_{th} [33]. Considering the threshold stress, eq. (1) can be rewritten as

$$\dot{\epsilon} = A \frac{Gb}{kT} \left(\frac{b}{d} \right)^p \left(\frac{\sigma - \sigma_{th}}{G} \right)^n D \quad (2)$$

The σ_{th} may be determined by adopting a plot of steady state stress σ against $\dot{\epsilon}^{1/n}$ ($n = 1, 2, 3, 5$) in a double linear scale [34]. The best linear fit in the entire temperature range was found for $n = 3$. By extrapolation to zero strain-rate (see Fig. 9) the threshold stress values were estimated for various temperatures and are introduced in Table 2.

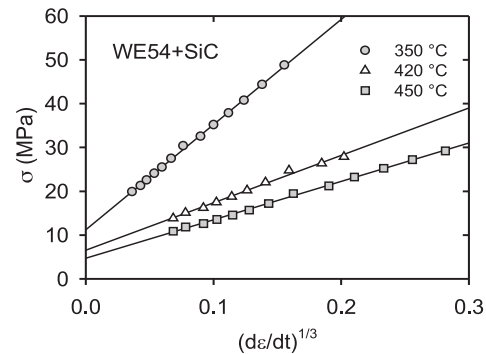


Fig. 9 Estimation of the threshold stress

From Table 2 it follows that the threshold stress decreases with increasing temperature. The temperature dependence of the normalised threshold stress may be expressed by an Arrhenius-type relationship in the form [35]

$$\frac{\sigma_{th}}{G} = B \exp \left(\frac{Q_0}{RT} \right) \quad (3)$$

Here B is constant and Q_0 is an energy term which appears to be associated with the deformation process. The activation energy Q_0 was estimated from the plot $\ln \dot{\epsilon} T/G$ vs $1/T$ (Fig. 10), measured at a constant normalized flow stress $(\sigma - \sigma_{th})/G$ ($\sigma - \sigma_{th}/G = 1.7 \times 10^{-3}$). The activation energy was found to be $Q \approx 114$ kJ/mol. The activation energy estimated for volume diffusion is 135 kJ/mol and for grain boundary diffusion 92 kJ/mol [36]. Comparing with the experimental value of 114 kJ/mol (using rule of mixture), we may conclude that the measured activation energy

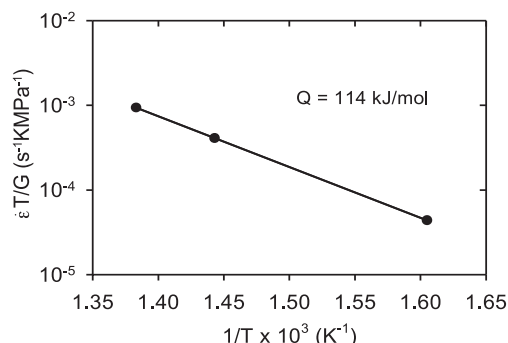


Fig. 10 Estimation of the activation energy

consists of approximately 50% volume and 50% grain boundary diffusion. The threshold stress is very probably necessary for the dislocation slip in the basal planes dependent on cross slip into non-basal slip planes (pseudo-Peierls mechanism), and/or grain boundary sliding.

Observed formation and growth of cavities relaxes the stress concentration caused at the particles on sliding grain boundaries. Cavities, created by vacancy clustering, may nucleate if the stress concentration is not relieved sufficiently rapidly. Local tensile stress caused by sliding at interfaces may be written in the form [4]

$$\sigma_{slid} = \frac{0.92kTd_p\dot{\epsilon}dV_f}{\Omega D_L \left(1 + 5 \frac{\delta D_{GB}}{d_p D_L}\right)}, \quad (4)$$

where d_p is the particle diameter, $\dot{\epsilon}$ is the strain rate, d is the grain size, D_L is lattice diffusion and D_{GB} the grain boundary diffusion coefficient, δ is grain boundary width, Ω is atomic volume and V_f is the volume fraction of particles. The insufficiently accommodated grain boundary sliding process is the reason for cavitation and early failure of samples

4. Conclusions

1. Magnesium alloys QE22 and EZ33 exhibit superplastic behaviour at 420 °C and the strain rates in the interval $1-3 \times 10^{-4} \text{ s}^{-1}$.
2. The grain size refinement in alloys may be attributed to very stable $\text{Mg}_3(\text{Ag}, \text{Nd})$ phase and T phase in grain boundaries formed in both materials during the preparation process.
3. Small addition of Zr particles significantly influences the resulting grain size.
4. SiC particles very probably inhibit the grain boundary sliding and they are reason for the premature failure of the WE54/SiC_p composite.
5. The threshold stress values found at temperatures 350 – 450 °C may be interpreted as the stresses necessary for the glide of dislocations in the compact and non-compact planes and grain boundary sliding.

Acknowledgements.

The work is a part of activities of the Charles University Research Center “Physics of Condensed Matter and Functional Materials”. The authors are grateful also for the financial support of the Czech Grant Agency under the contract P108/12/J018.

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