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Thandorn, T. and Tsakiropoulos, P. orcid.org/0000-0001-7548-3287 (2021) The effect of boron on the microstructure and properties of refractory metal intermetallic composites (RM(Nb)ICs) based on Nb-24Ti-xSi (x = 16, 17 or 18 at.%) with additions of Al, Cr or Mo. Materials, 14 (20). 6101. ISSN 1996-1944

https://doi.org/10.3390/ma14206101

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Article



The Effect of Boron on the Microstructure and Properties of Refractory Metal Intermetallic Composites (RM(Nb)ICs) Based on Nb-24Ti-xSi (x = 16, 17 or 18 at.%) with Additions of Al, Cr or Mo

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Abstract: This paper is about metallic ultra-high temperature materials, in particular, refractory metal intermetallic composites based on Nb, i.e., RM(Nb)ICs, with the addition of boron, which are compared with refractory metal high entropy alloys (RHEAs) or refractory metal complex concentrated alloys (RCCAs). We studied the effect of B addition on the density, macrosegregation, microstructure, hardness and oxidation of four RM(Nb)IC alloys, namely the alloys TT2, TT3, TT4 and TT8 with nominal compositions (at.%) Nb-24Ti-16Si-5Cr-7B, Nb-24Ti-16Si-5Al-7B, Nb-24Ti-18Si-5Al-7B, Nb-24Ti-18Si-5 5Cr-8B and Nb-24Ti-17Si-3.5Al-5Cr-6B-2Mo, respectively. The alloys made it possible to compare the effect of B addition on density, hardness or oxidation with that of Ge or Sn addition. The alloys were made using arc melting and their microstructures were characterised in the as cast and heat-treated conditions. The B macrosegregation was highest in TT8. The macrosegregation of Si or Ti increased with the addition of B and was lowest in TT8. The alloy TT8 had the lowest density of 6.41 g/cm^3 and the highest specific strength at room temperature, which was also higher than that of RCCAs and RHEAs. The Nbss and T2 silicide were stable in the alloys TT2 and TT3, whereas in TT4 and TT8 the stable phases were the Nb_{ss} and the T2 and $D8_8$ silicides. Compared with the Ge or Sn addition in the same reference alloy, the B and Ge addition was the least and most effective at 800 °C (i.e., in the pest regime), when no other RM was present in the alloy. Like Ge or Sn, the B addition in TT2, TT3 and TT4 did not suppress scale spallation at 1200 °C. Only the alloy TT8 did not pest and its scales did not spall off at 800 and 1200 °C. The macrosegregation of Si and Ti, the chemical composition of Nb_{ss} and T2, the microhardness of Nb_{ss} and the hardness of alloys, and the oxidation of the alloys at 800 and 1200 °C were also viewed from the perspective of the alloy design methodology NICE and relationships with the alloy or phase parameters VEC, δ and $\Delta \chi$. The trends of these parameters and the location of alloys and phases in parameter maps were found to be in agreement with NICE.

Keywords: Nb silicide-based alloys; high entropy alloys; complex concentrated alloys; refractory metal intermetallic composites; alloy design; intermetallics; solid solution; oxidation

1. Introduction

Research is underway worldwide to develop new metallic ultra-high temperature materials (UHTMs) to replace Ni superalloys in the hottest parts of future aero-engines to enable the latter to meet environmental and performance targets. Candidate metallic UHTMs include refractory metal (RM) intermetallic composites (RMICs), RM high entropy alloys (RHEAs) and RM complex concentrated alloys (RCCAs) (unnecessarily, in our opinion, another term has been added for RHEAs and RCCAs to further muddle the names used for metallic UHTMs, namely refractory chemically complex alloy (RCCAs)). The new materials must meet specific property targets (goals) for toughness, creep and oxidation



Citation: Thandorn, T.; Tsakiropoulos, P. The Effect of Boron on the Microstructure and Properties of Refractory Metal Intermetallic Composites (RM(Nb)ICs) Based on Nb-24Ti-xSi (x = 16, 17 or 18 at.%) with Additions of Al, Cr or Mo. *Materials* **2021**, *14*, 6101. http:// doi.org/10.3390/ma14206101

Academic Editor: Hideki Hosoda

Received: 21 August 2021 Accepted: 4 October 2021 Published: 15 October 2021

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Copyright: © 2021 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (https:// creativecommons.org/licenses/by/ 4.0/). resistance [1]. RMICs include materials based on Nb (RM(Nb)ICs, also known as Nb silicide-based alloys or Nb in situ composites) and Mo (RM(Mo)ICs). Recently, RM(Nb)ICs, RHEAs and RCCAs were compared in [1]. Some RM(Nb)ICs with B addition are also RCCAs [2] (i.e., RM(Nb)ICs/RCCAs). This paper is about RM(Nb)ICs with B addition and Ti/Si > 1.3 [1].

According to the alloy design methodology NICE (Niobium Intermetallic Composite Elaboration) [3], the alloying behaviour of RM(Nb)ICs and their phases, namely bcc Nb solid solution(s), M₅Si₃ silicides (M = RM = Mo, Ta or W, or M=TM (transition metal) = Cr, Hf or Ti), C14–NbCr₂ based Laves and A15-Nb₃X (X = Al, Ge, Si or Sn) compounds can be described using the parameters ΔH_{mix} , ΔS_{mix} , δ , VEC, $\Delta \chi$, Ω (=T_m $\Delta S_{mix} / |\Delta H_{mix}|$) that are also used to describe the alloying of HEAs (see Appendix A). RM(Nb)ICs and their phases, and HEAs can be presented in maps of the above parameters [4–9]. Solid solution(s) RHEAs and RCCAs, and multiphase RCCAs (bcc solid solution(s) + M₅Si₃ silicide with/without Laves phase) can be presented in maps of aforementioned parameters together with RM(Nb)ICs and their bcc solid solution(s) [1]. The RM(Nb)ICs with B addition occupy a separate distinct area in the maps [1,2,5].

Boron in RM(Nb)ICs partitions to the bcc solid solution, the 5-3 silicide [4,6,10,11] and Nb₃Si [11,12]. According to the Nb–Si–B phase equilibria, the 5-3 silicides are (a) the tetragonal T2 with Cr_5B_3 prototype, the same as $D8_1 \alpha Nb_5Si_3$, where B substitutes Si and becomes Nb₅(Si,B)₃, and (b) the hexagonal silicide with Mn₅Si₃ prototype, which is the same for the D8₈ γNb_5Si_3 , i.e., the metastable 5-3 silicide in the Nb–Si binary [13], and the D8₈ Ti₅Si₃ [14], and in which B occupies the interstitial site of Si and the silicide becomes Nb₅Si₃B_x [10]. The latter is referred to as the D8₈ silicide.

According to the 1600 °C isothermal section of Nb–Si–B by Nowotny et al. [10], the B and Si concentrations in the T2 and D8₈ silicides are in the ranges (at.%) 5.5 < B < 21 and 17 < Si < 33, and 5 < B < 16 and 27 < Si < 37.5, respectively, the T2 and D8₈ have homogeneity range along the direction parallel to the Si-B binary [10,15] and a composition variation (≤ 2.5 and 5.5 at.%, respectively) along the direction parallel to the Nb–Si binary [10] and the Nb content of the D8₈ is in the range 54 < Nb < 60. The D8₈ has Si + B \leq 42.5 at.% whereas the T2 has Si + B around 38 at.%. At 1600 °C, Sun et al. [11] calculated the maximum solubility of B in T2 to be 23.5 at.%, the B concentration in D8₈ in the range 10 < B < 15 at.% along the Si–B direction and the Nb concentration in D8₈ to be 55.6 \pm 1 at.%. The latter is in good agreement with the average of 57 at.% [10] and gives Nb/(Si + B) about 1.3 in D8₈ compared with 1.7 for the T2. The B solubility in the D8₈ decreases with increasing temperature [15].

In this paper, we study the microstructure and properties of four RM(Nb)ICs based on Nb-24Ti-xSi-yB with additions of Al, Cr or Mo (x = 16, 17 or 18, y = 6, 7 or 8) that were selected so that they belong in area C in the $\Delta\chi$ versus δ , ΔH_{mix} versus $\Delta\chi$, ΔH_{mix} versus VEC or VEC versus δ maps in [1,2,5]. The nominal composition of each alloy (at.%) was Nb-24Ti-16Si-5Cr-7B (alloy TT2), Nb-24Ti-16Si-5Al-7B (TT3), Nb-24Ti-18Si-5Al-5Cr-8B (TT4) and Nb-24Ti-17Si-3.5Al-5Cr-6B-2Mo (TT8), respectively (Table 1). These alloys are essentially based on specific KZ series alloys to which B has been added. In other words, the basis or reference alloys for TT2, TT3, TT4 and TT8, respectively are the alloys KZ4, KZ7, KZ5 [16] and JG3 [17] (for the nominal compositions of the reference alloys see Table A1 in Appendix A). Relevant data for the reference alloys are used to study how the addition of B in each alloy of this study affected microstructure and properties.

A 11 or 1		Element								
Alloy	Condition	Nb	Ti	Si	Al	Cr	В	Мо		
TT2	nominal	48	24	16	-	5	7	-		
-	AC	47.8	24.2	16.3	-	5.2	6.5	-		
-	HT	47.3	24	16.5	-	5.3	6.9	-		
TT3	nominal	48	24	16	5	-	7	-		
-	AC	49.3	23.6	15.4	4.4	-	7.3	-		
-	HT	49.5	23.2	15.6	4.4	-	7.3	-		
TT4	nominal	40	24	18	5	5	8	-		
-	AC	42.4	24.6	15.7	5	5.4	6.9	-		
-	HT	40.7	24	17.3	4.8	4.9	8.3	-		
TT8	nominal	42.5	24	17	3.5	5	6	2		
-	AC	43.4	23	17.2	3.5	4.9	6	2		
-	HT	42.9	23.7	17.2	3.3	4.9	5.9	2.1		

Table 1. Nominal and actual composition (at.%) of as cast (AC) and heat-treated (HT) alloys.

The structure of the paper is as follows: the description of experimental procedures is followed by the presentation of the results for the microstructures, density and hardness of the alloys. Preliminary results of the oxidation of the alloys are presented to show the effect of the addition of B on oxidation at 800 and 1200 °C. The discussion of the results includes their scrutiny from the perspective of NICE.

2. Experimental Section

Large button ingots (0.6 kg) of the alloys (Table 1) were prepared from high purity (min. 99.99 wt.%) Al, Cr, Mo, Nb, Si, Ti and B (99.5 wt.%) in an argon atmosphere using arc melting with a non-consumable tungsten electrode and a water-cooled copper crucible. Each alloy was melted five times. The alloys were heat-treated for 100 h at 1500 °C (1200 °C for TT2, see next section). Each alloy was wrapped in Ta foil and was heat-treated in a tube furnace under a continuous flow of Ti gettered argon followed by furnace cooling to room temperature. The microstructures of the alloys were characterised with XRD (Philips diffractometer, Hiltonbrooks Ltd., Crewe, UK, Cu radiation, solid specimens, JCPDS database) and microanalysis using EPMA (JEOL 8600 electron probe micro analyser, Tokyo, Japan, operating conditions 9 kV with a beam current of 30 nA [18]) equipped with WDX and EDX spectrometers, elemental standards and BN and TiN standards. The samples were prepared following standard metallographic procedures, mounted in Bakelite and ground using SiC grinding papers (320 to 2400 grit) and polishing with diamond DUR clothes (6 and 1 μ m finish).

The density of the alloys was measured using Archimedes principle and a Sartorious electronic precision balance, Sartorius Lab Instruments GmbH & Co. KG, Göttingen, Germany, equipped with a density measurement kit. The isothermal oxidation of the as cast (AC) alloys was studied at 800 and 1200 °C using Stanton Redcroft thermo-balances (Thermal Scientific plc., Odessa, TX, USA) and specimens from the as cast alloys. The weight of each sample was measured at the start and end of each oxidation experiment. A Mitutoyo HM-101 hardness testing machine (Mitutoyo America, Aurora, IL, USA) with Vickers indenter was used to measure the hardness (0.5 kg load) of the as cast (AC) and heat-treated (HT) alloys. The microhardness of the solid solution and T2 silicide was measured in the AC and HT alloys using a 0.025 kg load. At least ten measurements were done for each alloy and condition, and each phase. The area fraction of the solid solution was measured using the software KS Run 3 in a Polyvar Met microscope. The SEM images

of the large areas (\times 250) of the as cast (AC) and heat-treated (HT) alloys that were analysed by EPMA were used for the measurements.

3. Results

The actual composition of each alloy in the AC or HT condition is given in Table 1. The densities of the alloys are given in Table 2. The alloys TT4 and TT2 had the lowest and highest density, respectively, and the densities of all four alloys were lower than 7 g/cm³. The macrosegregation of solutes (MACX, X = Al, B, Cr, Si, Ti) in the alloys is given in Table 3. Note that this table has data for the Al and Cr free alloy TT1, to help us understand the role of boron with other alloying additions on MACX. The highest macrosegregation of B and Si was observed respectively in the alloys TT8 and TT4 whereas the macrosegregation of Ti was essentially similar in all four alloys. The phases that were observed in the alloys using XRD and EPMA are summarised in Table 4. The chemical compositions of the phases that were observed in all parts of the ingots are given in Table 5.

Table 2. Density of alloys, vol.% Nb_{ss}, hardness * of alloys, microhardness * of Nb_{ss} and T2 silicide.

	Hardness HV _{alloy}				XX 1 0/ XX1			Microhardness HV _{phase}			
Alloy			ρ (g/	ρ (g/cm ³)		V01.% Nb _{ss}		Nb _{ss}		T2	
	AC	HT	AC	HT	AC	HT	AC	HT	AC	HT	
TT2	$\begin{array}{c} 688 \pm 42 \\ 622 734 \end{array}$	$\begin{array}{c} 680 \pm 28 \\ 624 708 \end{array}$	6.71	6.72	37	38	$\begin{array}{c} 472\pm18\\ 443496\end{array}$	$\begin{array}{c} 419\pm39\\ 377495 \end{array}$	$\begin{array}{c} 1346 \pm 70 \\ 1202 1445 \end{array}$	$\begin{array}{c} 1167 \pm 84 \\ 1015 1265 \end{array}$	
TT3	$630 \pm 86 \\ 494 - 757$	$\begin{array}{c} 571 \pm 54 \\ 501 659 \end{array}$	6.52	6.56	38	42	$\begin{array}{c} 521 \pm 34 \\ 472 577 \end{array}$	$\begin{array}{c} 379 \pm 19 \\ 352 411 \end{array}$	$\begin{array}{c} 1232 \pm 72 \\ 1116 1334 \end{array}$	$\begin{array}{c} 1163 \pm 80 \\ 1061 1332 \end{array}$	
TT4	$\begin{array}{c} 720 \pm 70 \\ 642 879 \end{array}$	$\begin{array}{c} 607 \pm 21 \\ 590 659 \end{array}$	6.41	6.41	30	39	$\begin{array}{r} 563 \pm 37 \\ 503 631 \end{array}$	$\begin{array}{r} 467\pm15\\ 450496\end{array}$	$\begin{array}{c} 1247 \pm 55 \\ 1172 1334 \end{array}$	$\begin{array}{c} 1239 \pm 40 \\ 1200 1332 \end{array}$	
TT8	$744 \pm 60 \\ 673 - 842$	$693 \pm 28 \\ 622-717$	6.46	6.44	39	38	$\begin{array}{r} 531 \pm 26 \\ 487 576 \end{array}$	$\begin{array}{r} 461\pm25\\ 417495\end{array}$	$\begin{array}{c} 1316 \pm 37 \\ 1265 1369 \end{array}$	$\begin{array}{c} 1209 \pm 48 \\ 1143 1299 \end{array}$	

* given as the average value, standard deviation, and maximum and minimum measured values.

Table 3. Macrosegregation (at.%) of solutes MACX (X = Al, B, Cr, Si, Ti) in the cast alloys, where MACX = $C_{max}^{X} - C_{min}^{X}$ [19].

Alloy	MACX (at.%)									
	Al	В	Cr	Si	Ti					
TT2	-	4.2	2.8	3.1	3.2					
TT3	2.7	4.2	-	4.5	3.4					
TT4	1.1	2.9	1.4	4.7	3.3					
TT8	1.1	4.9	1.3	1.3	2.8					
TT1 [12]	-	4.4	-	4.1	5.5					

Table 4	Phases in	the as cast ((AC) and	heat-treated	(HT) all	oys accordin	g to th	e XRD	and EPMA	data.
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A 11		Phase												
Condition	Nb _{ss}	T2	Cr & Ti Rich Phase	Ti Rich Phase	C14-Laves	Nb ₃ Si	D8 ₈	Nb _{ss} + T2 Eutectic	TiN					
TT2-AC	X *	X *	Х	Х	Х	Х	-	Х	-					
TT2-HT [◊]	Х	X *	-	-	-	-	-	-	X °					
TT3-AC	X *	X *	-	-	-	-	-	Х	-					
TT3-HT +	Х	X *	-	-	-	-	-	-	-					
TT4-AC	X *	X *	-	-	-	-	Х	Х	-					
TT4-HT +	Х	Х	-	-	-	-	Х	-	-					
TT8-AC	X *	X *	-	-	-	Х	Х	Х	-					
TT8-HT +	Х	X *	-	-	-	-	Х	-	Χ°					

* indicates the presence also of Ti-rich phase (see Table 5), + 1500 °C/100 h, \diamond 1200 °C/100 h, ° indicates that alloy was contaminated by nitrogen.

Alloy and	Diana				Element			
Condition	rnase	Nb	Ti	Si	Al	В	Cr	Мо
TT2-AC	Nb _{ss}	$\begin{array}{c} 57.263.2 \\ 60.3 \pm 2.4 \end{array}$	$23.7-29 \\ 26.6 \pm 1.7$	$1.4-1.9 \\ 1.6 \pm 0.2$	-	$3.6-6.3 \\ 5 \pm 0.2$	5.3-7.9 6.5 ± 1	-
-	Ti rich Nb _{ss}	38.8-46.6 42.5 ± 3.8	36.9-43.9 40.7 ± 1.7	$\begin{array}{c} 1.63.5\\ 2.4\pm0.2\end{array}$	-	$0.6-3.3 \\ 1.8 \pm 0.2$	$\begin{array}{c} 10.114.1 \\ 12.6 \pm 2.3 \end{array}$	-
-	Phase rich in Cr & Ti	$\begin{array}{c} 14.821.9 \\ 18.9 \pm 2.6 \end{array}$	$\begin{array}{c} 40.253.9 \\ 48.3 \pm 1.4 \end{array}$	3.4-12.9 7.1 ± 2.7	-	-	$\begin{array}{c} 23.4 28.5 \\ 25.7 \pm 4.5 \end{array}$	-
-	T2	$46-48.1 \\ 47.1 \pm 0.6$	$\begin{array}{c} 13.4 {-} 15.4 \\ 14.3 \pm 0.9 \end{array}$	$\begin{array}{c} \textbf{23.4-26.4}\\ \textbf{25.3}\pm \textbf{1} \end{array}$	-	$\begin{array}{c} 10.9 {-} 14.9 \\ 12.9 \pm 1.1 \end{array}$	$0.2-0.7 \\ 0.4 \pm 0.1$	-
-	Ti rich T2	$\begin{array}{c} 39.1 43.7 \\ 41.6 \pm 1.2 \end{array}$	$\begin{array}{c} 18.6 23.8 \\ 20.8 \pm 1.6 \end{array}$	$\begin{array}{c} 28.1 30.3 \\ 29.3 \pm 0.7 \end{array}$	-	5.9-9.4 7.8 ± 1	$0.2-0.8 \\ 0.5 \pm 0.1$	-
TT3-AC	Nb _{ss}	52.9-59.4 55.8 ± 0.6	28.6-32.6 30.6 ± 1.2	$1.6-2 \\ 1.8 \pm 0.1$	3.9-8.4 6.8 ± 2.2	$\begin{array}{c} 46\\5\pm0.6\end{array}$	-	-
-	Ti rich Nb _{ss}	$\begin{array}{c} 43.449.9\\ 47.7\pm0.6\end{array}$	$\begin{array}{c} 38.7 - 47.8 \\ 42.7 \pm 1.6 \end{array}$	$\begin{array}{c} 1.72\\ 1.9\pm0.1\end{array}$	$\begin{array}{c} 3.35.4\\ 4.5\pm2.4\end{array}$	$\begin{array}{c} 1.94.4\\ 3.2\pm0.7\end{array}$	-	-
-	T2	$\begin{array}{c} 4648.3\\ 46.9\pm0.4\end{array}$	$\begin{array}{c} 13.315.6 \\ 14.4 \pm 0.2 \end{array}$	22.5-26.7 25 ± 1.2	$0.4-1.2 \\ 0.8 \pm 0.1$	$\begin{array}{c} 10.816.2 \\ 12.9 \pm 1.5 \end{array}$	-	-
-	Ti rich T2	$\begin{array}{c} 41.345.1\\ 42\pm1.4\end{array}$	$\begin{array}{c} 18.2 19.9 \\ 19 \pm 0.3 \end{array}$	$\begin{array}{c}\textbf{29.130.4}\\\textbf{29.8}\pm0.4\end{array}$	0.6-2.7 1.7 ± 0.6	$6.6-8.6 \\ 7.5 \pm 0.5$	-	-
TT4	Nb _{ss}	$46.4{-}50.1$ 48.3 ± 2.1	$\begin{array}{c} 29.4 31.6 \\ 30.6 \pm 1.7 \end{array}$	1.2-1.7 1.4 ± 0.1	$\begin{array}{c} \textbf{7.1-7.7}\\ \textbf{7.4} \pm \textbf{0.3} \end{array}$	2.5 - 3.9 3.3 ± 0.3	7.9-10.1 9 ± 0.4	-
-	Ti rich Nb _{ss}	$\begin{array}{c} 32.7 39.4 \\ 35.6 \pm 1.1 \end{array}$	$\begin{array}{c} 3742.5\\ 39.7\pm0.5\end{array}$	$\begin{array}{c} 1.11.5\\ 1.3\pm0.1\end{array}$	$\begin{array}{c} \textbf{6.7-7.3} \\ \textbf{7} \pm \textbf{0.4} \end{array}$	$\begin{array}{c} 0.31.2\\ 0.7\pm0.4\end{array}$	$\begin{array}{c} 14.116.9 \\ 15.7 \pm 1.4 \end{array}$	-
-	T2	$\begin{array}{c} 40.542.6\\ 41.4\pm0.3\end{array}$	$\begin{array}{c} 1920.3\\ 19.7\pm0.4\end{array}$	27.5-29.9 29.1 ± 0.8	$\begin{array}{c} 1.72.7\\ 2.3\pm0.3\end{array}$	$5.28.3$ 6.8 ± 0.4	$0.4{-}1.1$ 0.7 ± 0.1	-
-	Ti rich T2	28.5-35.8 32.5 ± 1	26.2-33.5 29.3 ± 0.9	$\begin{array}{c} \textbf{27.4-29.3} \\ \textbf{28.3} \pm \textbf{0.8} \end{array}$	$\begin{array}{c} 3.74.3\\ 4\pm0.3\end{array}$	$\begin{array}{c} 3.25.5\\ 4.4\pm0.4\end{array}$	0.8-2.2 1.5 ± 0.4	-
-	D8 ₈	$\begin{array}{c} 44.546.9\\ 45.9\pm0.6\end{array}$	$\begin{array}{c} 11.713.3 \\ 12.5\pm0.6 \end{array}$	$\begin{array}{c} 14.416\\ 15\pm0.8\end{array}$	-	$\begin{array}{c} 23.527\\ 25.6\pm0.4\end{array}$	$0.6-1.4 \\ 1 \pm 0.3$	-
TT8-AC	Nb _{ss}	$\begin{array}{c} 42.4 49.9 \\ 45.9 \pm 2.5 \end{array}$	$\begin{array}{c} 26.9 30.9 \\ 28.8 \pm 1.1 \end{array}$	$\begin{array}{c} 1.11.5 \\ 1.3 \pm 0.3 \end{array}$	$6.8-8.5 \\ 7.6 \pm 0.4$	$0-5.4 \\ 1.8 \pm 1.4$	$\begin{array}{c} 9.111.5 \\ 10.6 \pm 1.4 \end{array}$	$\begin{array}{c} 3.54.5\\ 4\pm0.6\end{array}$
-	Ti rich Nb _{ss}	$\begin{array}{c} 30.838\\ 34.5\pm0.7\end{array}$	36.4-41.5 38.4 ± 2.1	$\begin{array}{c}11.4\\1.2\pm0.3\end{array}$	$6-7.3 \\ 6.8 \pm 0.7$	$\begin{array}{c} 03.5\\ 1.1\pm0.8\end{array}$	$\begin{array}{c} 14.216.6 \\ 15.6 \pm 2.2 \end{array}$	$\begin{array}{c} \textbf{2-2.8}\\ \textbf{2.4}\pm \textbf{0.1} \end{array}$
-	T2	$\begin{array}{c} 36.7 43.5 \\ 40.5 \pm 1.2 \end{array}$	$\begin{array}{c} 18.220.8 \\ 19.1\pm0.4 \end{array}$	$\begin{array}{r} 2428\\ 25.6\pm4.9\end{array}$	$2.6 extrm{-}3.5$ 2.9 ± 0.2	$6.7 ext{}15.5$ 10.4 ± 6.3	$0.6-1.1 \\ 0.8 \pm 0.2$	$0.6-0.7 \\ 0.7 \pm 0.1$
-	Ti rich T2	$\begin{array}{c} 29.2 32.1 \\ 30.7 \pm 1.3 \end{array}$	$\begin{array}{c} 29.4 31.5 \\ 30.2 \pm 0.9 \end{array}$	$\begin{array}{c} 24.7 27.8 \\ 26.4 \pm 3.5 \end{array}$	3–4.2 3.3±0.5	$6.5-9.5 \\ 8 \pm 1.1$	$0.3-1 \\ 0.9 \pm 0.3$	$\begin{array}{c} 0.30.7 \\ 0.5 \pm 0.2 \end{array}$
-	D8 ₈	$\begin{array}{c} 41.945.2\\ 44.7\pm2\end{array}$	$\begin{array}{c} 9.212.3 \\ 10.2\pm0.6 \end{array}$	$\begin{array}{c} 1113.5 \\ 12.4\pm0.9 \end{array}$	-	26.4-33.7 30 ± 2.6	$0.8-1.3 \\ 1 \pm 0.2$	$\begin{array}{c} 1.61.8\\ 1.7\pm0.2\end{array}$
TT2-HT	Nb _{ss}	$52.1{-}55.2 \\ 53.2 \pm 1.2$	30.1-33.6 31.9 ± 1.6	$0.3-0.5 \\ 0.4 \pm 0.1$	-	$\begin{array}{c} 24.3\\ 3.2\pm0.5\end{array}$	10.2-12 11.3 ± 0.2	-
-	T2 *	46.9-48.6 47.8 ± 1.2	13.7-15.2 14.4 ± 1	23.3–26 24.5 ± 1.2	-	$10.7-14.8 \\ 12.7 \pm 1.5$	0.3-0.8 0.6 ± 0.1	-

 Table 5. EPMA data for the chemical composition (at.%) of phases in AC and HT alloys.

Alloy and	Dhaaa				Element			
Condition	Phase	Nb	Ti	Si	Al	В	Cr	Мо
TT3-HT	Nb _{ss}	54.8-57.9 56.4 ± 1	29.2-32 30.7 ± 1	$0.3-0.5 \\ 0.4 \pm 0.1$	7.5-8.3 7.8 ± 0.6	$\begin{array}{c} 3.85.6\\ 4.7\pm0.6\end{array}$	-	-
-	T2	$\begin{array}{c} 41.8 46.2 \\ 44.1 \pm 2.4 \end{array}$	$\begin{array}{c} 16.119.7 \\ 17.4 \pm 2.2 \end{array}$	$\begin{array}{c} 25.428\\ 26.9\pm2\end{array}$	$\begin{array}{c} 0.81.8\\ 1.4\pm0.9\end{array}$	$\begin{array}{c}911.7\\10.2\pm2.6\end{array}$	-	-
-	Ti rich T2	$34-36.6 \\ 34.8 \pm 1.3$	$\begin{array}{c} 2529\\ 27.4\pm1\end{array}$	$\begin{array}{c} 2829.5\\ 28.9\pm0.7\end{array}$	$\begin{array}{c} 3.74.5\\ 4.2\pm0.5\end{array}$	$\begin{array}{c} \textbf{3.8-5.9}\\ \textbf{4.7} \pm \textbf{0.4} \end{array}$	-	-
TT4-HT	Nb _{ss}	$\begin{array}{c} 48.5 50.5 \\ 49.8 \pm 0.8 \end{array}$	$\begin{array}{c} 28.1 29.5 \\ 28.6 \pm 0.6 \end{array}$	$\begin{array}{c} 0.40.7 \\ 0.5 \pm 0.1 \end{array}$	$\begin{array}{c} 7.37.8 \\ 7.6 \pm 0.2 \end{array}$	$\begin{array}{c} 1.54.3\\ 2.8\pm1\end{array}$	$\begin{array}{c} 10.411\\ 10.7\pm0.2\end{array}$	-
-	T2	$35.8{-}40$ 37.7 ± 1.9	$\begin{array}{c} 19.7 {-} 24.6 \\ 22.3 \pm 1.9 \end{array}$	$\begin{array}{c} 29.4 31.2 \\ 30.2 \pm 0.8 \end{array}$	$\begin{array}{c} 2.33.8 \\ 3.3 \pm 0.5 \end{array}$	4.2-6.3 5.5 ± 0.8	$0.7 ext{} 1.4 \\ 1 \pm 0.2$	-
-	D8 ₈	$\begin{array}{c} 42.7 - 44.5 \\ 43.6 \pm 0.8 \end{array}$	$\begin{array}{c} 11.612.4 \\ 11.9 \pm 0.3 \end{array}$	$\begin{array}{c} 12.9 14.2 \\ 13.4 \pm 0.5 \end{array}$	-	$\begin{array}{c} \textbf{28-30.7}\\ \textbf{29.2} \pm 1 \end{array}$	$\begin{array}{c} 1.62.2 \\ 1.9 \pm 0.3 \end{array}$	-
TT8-HT	Nb _{ss}	$\begin{array}{c} 46.2 - 48.7 \\ 45.6 \pm 1 \end{array}$	$\begin{array}{c} 23.9 - 31.5 \\ 29.8 \pm 2.2 \end{array}$	$\begin{array}{c} 0.40.6 \\ 0.5 \pm 0.1 \end{array}$	$\begin{array}{c} 7.28.2 \\ 7.6 \pm 0.5 \end{array}$	$\begin{array}{c} 04.8\\ 1\pm0.8\end{array}$	$\begin{array}{c} 10.511.8 \\ 11.2\pm0.7 \end{array}$	$\begin{array}{c} 4.14.4\\ 4.3\pm0.1\end{array}$
-	T2	$35.8-44.5 \\ 41 \pm 4.6$	$\begin{array}{c} 18.7 21.5 \\ 20.1 \pm 1.4 \end{array}$	$\begin{array}{c} 27.1 33.2 \\ 29.3 \pm 3.4 \end{array}$	$\begin{array}{c} 2.74.3 \\ 3.3 \pm 0.8 \end{array}$	$\begin{array}{c} 08.7\\ 4.7\pm4.4\end{array}$	$\begin{array}{c} 0.41.9 \\ 1.1 \pm 0.8 \end{array}$	$\begin{array}{c} 0.30.7 \\ 0.5 \pm 0.2 \end{array}$
-	Ti rich T2	$\begin{array}{c} 30.9 33.7 \\ 31.6 \pm 1.2 \end{array}$	$\begin{array}{c} 25.9 - 31.2 \\ 28.1 \pm 2 \end{array}$	26.4-29.6 28 ± 1.3	$\begin{array}{c} 3.33.6\\ 3.5\pm0.2\end{array}$	$\begin{array}{c} \textbf{3.3-11}\\ \textbf{6.8} \pm \textbf{4.6} \end{array}$	$\begin{array}{c} 1.11.8\\ 1.5\pm0.3\end{array}$	$\begin{array}{c} 0.40.5 \\ 0.5\pm 0.02 \end{array}$
-	D8 ₈	${41.2-48.9}\\ 44.2 \pm 4.3$	$\frac{11.412.6}{12.1\pm0.7}$	$\overline{ 12.44.9 } \\ 13.5 \pm 1.5 \\$	-	$ 19.2-31 \\ 26.5 \pm 6.4 $	$1.5-2.3 \\ 2 \pm 0.5$	1.6-1.9 1.7 ± 0.1

Table 5. Cont.

* see text.

3.1. As Cast Alloys

In all parts of the ingot of the alloy TT2, were observed the Nb_{ss}, T2, a Cr and Tirich phase and a eutectic of the Nb_{ss} and T2, whereas in specific parts of the ingot other phases were also present, namely a Ti-rich phase in the top and bottom, a C14-Laves phase in the top and bulk and Nb₃Si silicide in the bulk (Tables 4 and 5, Figure 1a and Figure S1a in Supplementary Materials). Ti-rich Nb_{ss} and Ti-rich T2 were also observed. In Figure 1a, the grey contrast facetted phase is the T2, the bright contrast phase is the Nbss, darker contrast areas at the edges of Nbss are Ti-rich Nbss and darker grey contrast areas in between Nb_{ss} are Ti-rich T2. The Ti-rich phase, C14-Laves phase, Cr and Ti-rich phase and Nb₃Si were observed in the dark and very contrast inter-dendritic Nb_{ss} areas. The average compositions of the Ti-rich phase, the Laves phase and the Nb₃Si silicide, respectively, were 23.2(18.6–27.9)Nb-52.3(47.8–56.7)Ti-8.9(8.4–9.4)Si-14.7(14–15.5)Cr-0.9(0.8– 1)B, 13.4(12.6–14.1)Nb-31.6(27.5–34.2)Ti-7.8(6.7–9)Si-47.7(45.3–49)Cr and 25.9(21.7–29)Nb-42.4(43-44.1)Ti-23.4(19.4-26.5)Si-7.9(4-11.3)Cr, where in the parentheses are given the minimum and maximum analysis values for each phase. It should be noted that the Laves and Nb₃Si compounds were B free and that the Si concentration in the silicide would suggest also the presence of metastable Nb_3Si [11], which however was not indicated by the XRD (Figure S1a in Supplementary Materials). In the Nb_{ss}, with increasing Ti concentration, the concentrations of B and Cr respectively decreased and increased, whereas in the T2 the B concentration decreased with increasing Ti concentration (Table 5). The average <Si> content and the <Nb>/<Si> and Si/B ratios of the T2 and Ti-rich T2 were 38.2 and 37.1 at.%, 1.6 and 1.7, and 2 and 3.8, respectively (where $\langle Nb \rangle = Nb + Cr + Ti$ and $\langle Si \rangle = B + Si$). The T2 silicide was facetted, this was particularly noticeable in the top and bottom of the ingot. The vol.% of the eutectic was lower in the bulk.



Figure 1. Microstructures of the AC (**a**,**c**,**e**,**g**) and HT (**b**,**d**,**f**,**h**) alloys TT2 (**a**,**b**), TT3 (**c**,**d**), TT4 (**e**,**f**) and TT8 (**g**,**h**). Note that contrast enhancement has been applied to show different phases. For description of microstructures see text.

In all parts of the ingot, the microstructure of the alloy TT3 consisted of the Nb_{ss}, T2, and eutectic of these two phases (Tables 4 and 5, Figure 1c and Figure S1c in Supplementary Materials). The XRD suggested the presence of the D8₈ silicide, however, its presence was not confirmed by EPMA. Ti-rich Nb_{ss} and T2 were also observed but in this alloy the partitioning of Ti to T2 was very strong and resulted in distinct separate T2 grains very rich in Ti (and Al) in the top and bulk of the ingot, with average composition 34.2(29.8–37.2)Nb-28.2(24.7–32.2)Ti-27.8(26.6–28.8)Si-4.2(3.7–4.9)Al-5.6(3.6–7.4)B. In Figure 1c, the light contrast phase is the Nb_{ss}, the grey contrast phase is the T2 and the darker grey contrast is the Ti-rich T2. In the Nb_{ss} and T2, with increasing Ti concentration, the concentrations of both B and Al decreased in the solid solution, and respectively decreased and increased in the T2. The average <Si> content, and the <Nb>/<Si> and Si/B ratios of the T2, Ti-rich T2 and T2 very rich in Ti were 38.6, 39 and 37.5 at.%, 1.6, 1.56 and 1.67, and 1.9, 4 and 5, respectively (where <Nb> = Nb + Ti and <Si> = Al + B + Si). The T2 was facetted only in the bulk of the ingot. Cracks that had formed in large Ti-rich T2 grains were stopped by the surrounding solid solution.

In all parts of the ingot of the alloy TT4, the Nb_{ss}, T2, D8₈ and eutectic of Nb_{ss} and T2 were observed. The D8₈ silicide exhibited very bright contrast (Tables 4 and 5, Figure 1e and Figure S1e in Supplementary Materials). Ti-rich Nb_{ss} and Ti-rich T2 were also observed but there was no evidence of Ti-rich D8₈. The latter was Al free. In Figure 1e, the T2 is the grey contrast phase, darker grey contrast areas correspond to Ti-rich T2, the very bright contrast thin long phase at the bottom left-hand side, middle and upper right-hand side of the image "running" at about 45 degrees from right to left) and the bright phase is the Nb_{ss} with darker areas corresponding to the Ti-rich Nb_{ss}.

In the Nb_{ss} and T2, with increasing Ti concentration, the concentrations of B and Cr decreased and increased, respectively, and the concentration of Al essentially did not change in the solid solution, whereas in the T2 the Al and Cr increased and the B decreased. The average $\langle Si \rangle$ content, and the $\langle Nb \rangle / \langle Si \rangle$ and Si/B ratios of the T2, Ti-rich T2 and D8₈ were 38.2, 36.7 and 40.8 at.%, 1.62, 1.72 and 1.45, and 4.3, 6.4 and 0.6, respectively (where $\langle Nb \rangle = Nb + Ti + Cr$ and $\langle Si \rangle = Al + B + Si$). The T2 was facetted in the top and the bulk of the ingot and Ti-rich T2 grains were cracked with the cracks parallel to each other and not extending in the surrounding Nb_{ss}.

The microstructure of the alloy TT8 consisted of Nb_{ss} , T2, D8₈, Nb_{ss} + T2 eutectic in all parts of the ingot and Nb₃Si that was observed only in the top and bulk of the ingot (Tables 4 and 5, Figure 1g and Figure S1g in Supplementary Materials). The average composition of Nb₃Si was 46Nb-23Ti-20.1Si-2.8Al-10Cr-5.8B-1.3Mo. The D8₈ exhibited very bright contrast and was Al free. In Figure 1g, the T2 is the grey contrast phase, darker grey contrast areas correspond to Ti-rich T2, the very bright contrast phase with no darker contrast areas around it is the D8₈, and the bright phase with darker areas is the Nb_{ss}. The Nb₃Si exhibits very dark contrast in the inter-dendritic Nb_{ss} areas. The concentration of B in the solid solution varied significantly. It should be noted that there were solid solution grains that were B free. Ti-rich Nbss and Ti-rich T2 were also observed, and no Ti-rich D8₈. In the Nb_{ss} and T2, with increasing Ti concentration, the concentrations of Al, B and Mo decreased and that of Cr increased in the solid solution whereas in the T2 the B decreased but the changes of Al, Cr and Mo were minimal. The average <Si> content, and the $\langle Nb \rangle / \langle Si \rangle$ and Si/B ratios of the T2, Ti-rich T2 and D8₈ were 38.9, 37.7 and 42.6 at.%, 1.57, 1.65 and 1.35, and 2.5, 3.3 and 0.4, respectively (where <Nb> = Nb + Ti + Cr and $\langle Si \rangle = Al + B + Si \rangle$. Both the T2 and D8₈ were facetted, particularly the latter and there were cracks in T2 that run parallel to each other and stopped at the Nb_{ss}.

The vol.% of the Nb_{ss} was essentially the same in the alloys TT2, TT3 and TT8, about 38% and significantly lower in the alloy TT4 (Table 2). The microhardness of the solid solution was lowest and highest respectively in the alloys TT2 and TT4 (472 and 563 HV, respectively), the microhardness of T2 was highest and lowest in the alloys TT2 and TT3, respectively (1346 and 1232 HV, respectively), and the highest and lowest alloy hardness

was measured for the alloys TT8 and TT3, respectively (744 and 630 HV, respectively), see Table 2.

3.2. Heat Treated Alloys

All the alloys were heat-treated at 1500 °C with the exception of the alloy TT2 that exhibited liquation at this temperature and was heat-treated at 1200 °C. The phases that were present in their microstructures are given in Table 4. The alloys TT2 and TT8 were contaminated by nitrogen during the heat treatment.

In the alloy TT2 the Nb₃Si silicide, the Ti-rich phase, the Cr and Ti-rich phase and the Laves phase that were present in the cast alloy were not observed. The Nb_{ss} and T2 were present with Ti-rich areas in the latter (Tables 4 and 5, Figure 1b and Figure S1b in Supplementary Materials). In Figure 1b, the light contrast areas are the Nb_{ss}, the grey contrast phase is the T2, the darker grey contrast is the Ti-rich T2 and the very dark areas are TiN. The chemical inhomogeneity that was characteristic of this alloy in the as cast condition resulted in T2 grains in the heat-treated microstructure with <Nb>/<Si> and Si/B ratios covering a wider range than in the other alloys. The chemical composition of T2 and Ti-rich T2 did not change significantly compared with the cast alloy (Table 5), the former had $\langle Si \rangle = 37.2$ at.% ($\langle Nb \rangle / \langle Si \rangle = 1.69$) and Si/B about 1.9 whereas the latter had Si + B in the range 34.1 to 37 at.% (or $\langle Nb \rangle / \langle Si \rangle$ in the range 1.7 to 1.93) and Si/B about one. The average composition of the Ti-rich T2 was as follows in different parts of the heat-treated ingot: in the top 39.9(37.7-42.4)Nb-20.8(17.4-24.1)Ti-18.6(17.1-19.6)Si-2.2(1.8-2.9)Cr-18.5(16.7-20.2)B, in the bulk 42.4(41.4-43.9)Nb-20.6(19.4-21.2)Ti-19.7(18.4-21.4)Si-2.2(1.7–2.7)Cr-15.2(13.3–17.6)B, and in the bottom (where only one analysis was possible) 41.5Nb-19.2Ti-18.1Si-2.4Cr-18.9B. In the bottom of the heat-treated alloy there were also areas of the Ti-rich T2 where the Si and B concentrations were not equal, with $\langle Si \rangle = 37$ at.% (<Nb>/<Si> = 1.7) and Si/B about 3.8 (41.7(39.6–43.5)Nb-20.8(18.4–22.4)Ti-29.3(27.9– 30.3)Si-7.7(6.5–9.8)B-0.5(0.4–0.7)Cr). Furthermore, there were Ti-rich T2 grains with $\langle Si \rangle =$ 35.1 at.% (<Nb>/<Si> = 1.85) and Si/B = 6.3 (where <math><Si> = Si + B, <Nb> = Nb + Cr + Ti).

The microstructures of the alloys TT3 and TT4 consisted of the Nb_{ss}, T2 with the D8₈ present only in the latter alloy and no TiN in both alloys. Ti-rich T2 was observed in TT3 but not in TT4 and the D8₈ was Al free, as was the case in the cast alloy (Tables 4 and 5, Figure 1d,f and Figure S1d,f in Supplementary Materials). The microstructure of TT3-HT is shown in Figure 1d, where the light contrast phase is the Nb_{ss} , the grey contrast phase is the T2 and the darker grey contrast is the Ti-rich T2. In the alloy TT3, the average <Si> content, and the <Nb>/<Si> and Si/B ratios of the T2 and Ti-rich T2 were 38.5 and 37.7 at.%, 1.6 and 1.65, and 2.6 and 6.1, respectively (where $\langle Nb \rangle = Nb + Ti$ and $\langle Si \rangle = Al + Ti$ B + Si). In the TT4-HT there were no facetted T2 grains and often the D8₈ silicide was observed inside (surrounded by) T2 grains. The microstructure of TT4-HT is shown in Figure 1f, where the bright phase is the Nb_{ss} , the grey phase is the $D8_8$ and the darker grey contrast phase is the T2. The T2 had $\langle Si \rangle = 39$ at.%, $\langle Nb \rangle / \langle Si \rangle = 1.56$, Si/B = 5.5 ($\langle Si \rangle = 1.56$) Al + B + Si, $\langle Nb \rangle = Nb + Cr + Ti$, whereas the $D8_8$ had $\langle Si \rangle = 42.7$ at.%, $\langle Nb \rangle / \langle Si \rangle = 42.7$ 1.3 and Si/B = 0.5. In the alloy TT8 the microstructure consisted of Nb_{ss}, the T2, D8₈ and Ti-rich T2 silicides, was contaminated by nitrogen and the Nb₃Si was not present (Tables 4and 5, Figure 1h and Figure S1h in Supplementary Materials). The D8₈ was Al free. The microstructure of TT8-HT is shown in Figure 1h, where the bright phase is the Nb_{ss} , the grey phase is the D8₈, the darker grey contrast phase is the T2 and dark areas around the latter correspond to Ti-rich T2. The average <Si> content, and the <Nb>/<Si> and Si/B ratios of the T2, Ti-rich T2 and D8₈ were 37.3, 38.3 and 40 at.%, 1.68, 1.61 and 1.5, and 6.2, 4.1 and 0.51, respectively ($\langle Nb \rangle = Nb + Cr + Mo + Ti$). Some grains for the Nb_{ss} were B free, as in TT8-AC.

After the heat treatment, the vol.% of Nb_{ss} increased slightly in the alloys TT2, TT3 and TT8 and significantly in TT4 and was similar in all alloys, about 39% (Table 2). The microhardness of the Nb_{ss} was reduced in all alloys, most noticeably in TT3. Similarly, the microhardness of T2 was reduced in all alloys, but the smallest reduction was in TT4. The

hardness of all alloys was reduced, the TT8 still had the highest hardness but the smallest hardness reduction was measured for TT2 (Table 2).

3.3. Oxidation

The mass change data is shown in Figure 2a,b for isothermal oxidation at 800 and 1200 °C and typical examples of scale spallation at the two temperatures are shown in Figure 2c,d. The Figure 2a,b include data for the MASC alloy (Nb-25Ti-16Si-2Al-2Cr-8Hf [20]) and the alloy TT1 [12]. The oxidation rate constants are given in Table 6. At 800 and 1200 °C, all four alloys followed respectively linear and parabolic oxidation kinetics. None of the alloys with B addition suffered from pest oxidation at 800 °C, meaning each alloy did not convert into powder, but the scales that formed on the alloys TT1, TT2, TT3 and TT4 spalled off. At 1200 °C scale spallation was also observed for the alloys TT1, TT2, TT3 and TT4, and the MASC alloy. The 1200 °C specimen of TT8 looked exactly like the specimen shown in Figure 2d.

Table 6. Oxidation rate constants and mass changes of the alloys after isothermal oxidation for 100 h at 800 and 1200 °C.

Allov _	Oxidation R	ate Constant	Mass C ΔW/A (m	hange 1g/cm²)	Pest Oxidation	Scale Sj	pallation
intoy	800 °C k _l (g/cm ² s)	1200 °C k _p (g²/cm ⁴ s)	800 °C	1200 °C	800 °C	800 °C	1200 °C
MASC	$1.53 imes 10^{-7}$	$1.58 imes 10^{-8}$	36 (≤65 h)	80	MC *	-	Yes
TT1	$1.86 imes 10^{-7}$	$1.53 imes 10^{-8}$	63	80.3	No	Yes	Yes
TT2	$1.89 imes 10^{-7}$	$3.32 imes 10^{-8}$	61.4	105.6	No	Yes	Yes
TT3	$1.22 imes 10^{-7}$	$1.21 imes 10^{-8}$	38.6	66.1	No	Yes	Yes
TT4	$8.9 imes10^{-8}$	$3.4 imes10^{-9}$	27.9	34	No	Yes	Yes
TT8	$3 imes 10^{-9}$	$3.3 imes10^{-9}$	1.4	30	No	No	No

* MC = Maltese cross.



Figure 2. Cont.



Figure 2. Mass change (Δ W/A) versus time of alloys TT1, TT2, TT3, TTT4, TT8 and MASC at (**a**) 800 °C and (**b**) 1200 °C. Colours: black TT1, blue TT2, green TT3, red TT4, yellow MASC, brown TT8. Specimens of TT4 and TT8 after oxidation (**c**) at 1200 °C and (**d**) 800 °C, respectively.

4. Discussion

4.1. Macrosegregation

The addition of B increased the macrosegregation of Ti and Si compared with the reference B free alloys, with the exception of TT8 where MACSi decreased compared with JG3 (Table 7). TT4 had the highest MACSi, an increase of 3.4 at.% compared with KZ5, and TT1 had the highest MACTi. The highest and lowest MACB was exhibited respectively by TT8 and TT4. With the exception of the alloy TT4, the spread of MACB was narrow and around 4.4 at.%.

Alloy	MACS	Si (at.%)	Alloy	M (a	MACB (at.%)		M (a	MACTi (at.%)	
TT4	•	4.7	TT8	A	4.9	TT1	A	5.5	
TT3	100.0755	4.5	TT1		4.4	TT3		3.4	
TT1		4.1	TT2		4.2	TT4		3.3	
TT2		3.1	TT3		4.2	TT2		3.2	
JG3		2.6	TT4		2.9	KZ3		3.1	
KZ7		2.3	-		-	TT8		2.8	
KZ4		1.9	-		-	KZ7		2.3	
KZ3		1.6	-		-	JG3		1.5	
TT8		1.3	-		-	KZ4		1.4	
KZ5		1.3	-		-	KZ5		1.4	

Table 7. Comparison of the macrosegregation (MACX) of X = B, Si or Ti in the B containing alloys of this work and their basis/reference alloys KZ3, KZ4, KZ5, KZ7 [16] and JG3 [17] (see Appendix A). Large button ingots (0.6 kg) of all alloys were prepared using arc melting. Note that the table includes data for the alloy TT1 (Nb-24Ti-18Si-8B [12]) and its basis alloy KZ3 (Nb-24Ti-18Si [16]).

The macrosegregation of Si and Ti in the alloys of this study and their reference alloys is shown in Figure 3a–p. This figure shows plots of MACSi and MACTi versus specific parameters that describe macrosegregation in RM(Nb)ICs [19]. In agreement with previous research on B free RM(Nb)ICs [21–24], RM(Nb)ICs/RCCAs [25] and HEAs [26] both MACSi and MACTi increased with ΔH_m^{sp} , $\Delta H_m/T_m$, $[\Delta H_m/T_m] \times [\Delta H_m^{sd}/\Delta H_m^{sp}]^{-1}$ and T_m^{sp} , and decreased with ΔH_m^{sd} , $\Delta H_m^{sd}/\Delta H_m^{sp}$, T_m^{sd} and T_m^{sd}/T_m^{sp} (for definitions and equations for above parameters, please refer to [19]). This means that alloy design in NICE can optimise both MACSi and MACTi in B containing alloys using the same parameters. It should be noted that in the MACSi plots the data for the alloy TT8 and its basis JG3 does not follow the aforementioned trend. Further research is essential to confirm MACSi in TT8 and to study RC(Nb)ICs based on TT4 with RM addition(s) as both alloys (i.e., TT4 and TT8) point to a "pathway" for developing RCCAs with B addition [1,2].



Figure 3. Cont.



Figure 3. Cont.



Figure 3. Cont.



Figure 3. Cont.



Figure 3. Cont.



Figure 3. Cont.



Figure 3. MACSi (**a**,**c**,**e**,**g**,**i**,**k**,**m**,**o**), MACTi (**b**,**d**,**f**,**h**,**j**,**l**,**n**,**p**) and MACB (**q**,**r**,**s**) versus parameters for alloys TT1, TT2, TT3, TT4 and TT8, and their basis/reference alloys KZ3, KZ4, KZ5, KZ7 and JG3. Diamonds for B containing alloys. Colours (**a**–**p**), brown TT1, KZ3, green TT2, KZ4, red TT3, KZ7, purple TT4, KZ5, blue TT8, JG3. Pink for the MACB data. The alloy TT8 is indicated in (**q**) to (**s**). If TT8 were to be excluded, the R² values for the parabolic fit of the data change to 0.9986, 0.813 and 1, respectively in (**q**,**r**,**s**).

The data for MACB is limited only to the alloys of this study and the alloy TT1 [12]. Figure 3q to s show a good parabolic fit of this limited data of MACB versus the parameters $\Delta H_m{}^{sd}$, $\Delta H_m{}^{sd}/\Delta H_m{}^{sp}$ and $T_m{}^{sd}$, and suggests "maximum" MACB between five and six at.% for $\Delta H_m{}^{sd} \approx 18 \text{ kJ/mol}$, $\Delta H_m{}^{sd} \approx 1.535 \text{ kJ/mol}$ and $T_m{}^{sd} \approx 1840 \text{ K}$. More data about MACB is needed before we can firmly ascertain that the same parameters and trends also describe the macrosegregation of B in RM(Nb)ICs, and RHEAs or RCCAs with B addition.

4.2. Microstructures

The alloy microstructures were chemically inhomogeneous, particularly in the case of the alloy TT2. In the reference alloys KZ7, KZ5 and JG3, the high-temperature tetragonal β Nb₅Si₃ had formed from the melt [16,17], whereas with the addition of B, in the alloys TT2, TT3, TT4 and TT8 the tetragonal T2 silicide (same structure as the low-temperature α Nb₅Si₃, see introduction) formed from the melt as did the hexagonal D8₈ silicide in the alloys TT4 and TT8. In the reference alloy KZ4, the α Nb₅Si₃ formed via the eutectoid transformation of Nb₃Si [16]. Solute partitioning made the identification of phases, in particular the D8₈ silicide in the alloys TT4 and TT8, difficult. In all four alloys, the Nb_{ss} and T2 were stable phases and the D8₈ in TT4 and TT8.

In the alloy TT2 the Ti-rich phase, Nb₃Si and C14 Laves phase formed in competition with the Cr and Ti-rich phase (Table 4) in specific parts of the ingot of the cast alloy. The Ti-rich phase was observed only in the top and bottom of the ingot, the Laves phase in the top and bulk, similarly to KZ4-AC [16], and the Nb₃Si silicide only in the bulk. The aforementioned four phases were not stable in the heat-treated alloy. Note that the C14 Laves phase was also not stable in KZ4-HT. The $Nb_{ss} + Nb_3Si$ eutectic that formed in KZ4-AC was replaced by the Nb_{ss} + T2 eutectic in TT2-AC. In the latter, the area fraction of the eutectic was significantly reduced in the bulk of the ingot rather than at the bottom, which was the case in TT1-AC [12], and would suggest sensitivity of the Nb_{ss} + T2 eutectic to cooling rate and the presence or not of the Nb₃Si silicide. The Laves phase formed from Cr rich melt and thus depended on the partitioning of Cr during solidification. This would explain why the Laves phase was not observed in the bottom of the ingot. Compared with KZ4-AC, no Ti-rich areas were observed in the Laves phase in TT2-AC and the Ti concentration was significantly higher than in the Laves phase in KZ4-AC. This was attributed to the Laves phase "competing for Ti" with the Cr and Ti-rich phase (which formed everywhere in the ingot) and the Ti-rich phase (which formed in the top and bottom of the ingot). Chemical inhomogeneity existed in TT2-HT, with a range of composition of the Ti-rich T2 (see Table A2 in the Appendix A) indicating that equilibrium was not reached in this alloy after 100 h at 1200 °C.

The as solidified microstructure in TT3 was similar to that in KZ7-AC [16], the Nb_{ss} + β Nb₅Si₃ eutectic in the latter was replaced by the Nb_{ss} + T2 eutectic in TT3-AC. Compared with TT1-AC, the addition of Al in TT3 suppressed the formation of Nb₃Si, which was also the case in KZ7-AC compared with KZ3-AC [16]. The suppression of the Nb₃Si was also linked with the strong partitioning of Ti in the T2 that "starved" the melt of Ti, which stabilises the Nb₃Si. Therefore, the synergy of B and Al in TT3 furthered the formation of primary tetragonal T2 (as did the addition of Al in KZ7 regarding the formation of primary β Nb₅Si₃ [16]) and of the Nb_{ss} + T2 eutectic. The partitioning of Ti in T2 was sensitive to cooling rate as the very Ti-rich T2 was not observed in the bottom of the ingot of TT3-AC.

In the alloy TT4, the C14 Laves phase was not formed, differently with KZ5-AC [16]. In the alloy TT8-AC, the Nb₃Si silicide was formed but not the C14 Laves, similarly with JG3-AC [17], in which Nb_{ss} + Nb₃Si eutectic was formed compared with the Nb_{ss} + T2 eutectic in TT8-AC. The Nb₃Si was not observed in the bottom of the ingot of TT8-AC. Comparison with the data for JG3-AC [17] and TT8-AC would suggest sensitivity of the formation of Nb₃Si to cooling rate that was attributed to the synergy of B, Cr and Mo in which B and Cr lead. Three-phase equilibria between Nb_{ss}, Nb₅Si₃ and Nb₄Si (metastable silicide) would exist for an alloy of composition Nb-17.5Si-2Mo (at.%) according to the ternary Nb-Si-Mo system by Savitskiym et al. [27]. The Nb₃Si in TT8-AC had Si + B + Al \approx 28.7 at.%, which is close to the composition of the metastable Nb₃Si-m in [13], and the Si content (20.1 at.%) corresponds to Nb₄Si (metastable silicide, [13]). Thus, it is likely that with the addition of Mo both the stable (i.e., tetragonal) and metastable Nb₃Si silicide(s) formed in TT8-AC. Compared with JG3-AC, the Nb₃Si was leaner in Si and Al and richer in Cr; this was attributed to the synergy of Mo and B in TT8. The concentration of Mo was the same in the Nb₃Si in JG3-AC and TT8-AC, which would suggest that B did not affect the concentration of Mo in the 3-1 silicide. Compared with TT1-AC [12], the Nb₃Si in TT8-AC was leaner in Si but richer in B. Compared with TT2-AC, the Nb₃Si was B free, it was concluded that in the presence of Mo the solubility of B in Nb₃Si is increased and that Mo leads in the Mo and Cr synergy and eliminates the negative effect of Cr on the solubility of B in Nb₃Si.

In both the TT4 and TT8 alloys (Ti/Si = 1.5 based on nominal compositions or average Ti/Si = 1.42 based on actual chemical compositions, Table 1), with the addition of B, a new B rich phase formed and was stable, namely the D8₈ silicide. The latter was observed in the areas surrounding the T2 and exhibited very strong bright contrast, which was stronger than that of Nb_{ss} (Figure 1e,g). There was no solubility of Al in D8₈ and its Ti concentration was in the range 10.2 < Ti < 12.5 at.% (average 11.7 at.%, Table 5). In all the alloys of this study, whether in the AC or HT condition, the Si + B or the Si + B + Al content of the T2 was essentially the same (37 and 38.1 at.%, respectively, average 37.6 at.%) whereas the Si + B content of the D8₈ was higher and in the range 1 to 6.4 (average 4) compared with 0.5 for the D8₈ silicide (range 0.4 to 0.6), and the average <Nb>/<Si> ratio was 1.7 in the T2 compared with 1.4 in the D8₈ (Table A2 in Appendix A). The Ti concentration of the T2 and D8₈, respectively, was in the ranges 14.3 to 30.7 at.% (average 21.2 at.%) and 10.2 to 12.5 at.% (average 11.7 at.%), Table 5. The above data for the T2 and D8₈ silicides is in agreement with the Nb-Si-B ternary [10], see introduction.

In the 1600 °C ternary Nb-Si-B section, tentative equilibrium between T2 and Nb₃B₂ was indicated by Nowotny et al. [10]. Thermodynamic modelling of the Nb–Si–B ternary [28] calculated the Nb₃B₂ to be stable at 1600 °C and showed that it formed two three-phase equilibria Nb + T2 + Nb₃B₂ and T2 + Nb₃B₂ + NbB with neighbouring phases. Sun et al. [28] stated "there are no direct experimental evidences to support the existence of these two three-phase equilibria" and "further experimental results are needed to accurately define the stability region of Nb₃B₂".

The Nb_3B_2 compound was reported by the same researchers [11] in the B-rich, arc melted and heat-treated (1500 °C/100 h) very small buttons (0.006 kg) of seven Nb–Ti–Si–B alloys, of which six were B-rich (B \geq 10 at.%) and one B-lean (B = 1.5 at.%), all alloys had 16 at.%Si, four had Ti/Si < 1 (range 0.28 to 0.9, Ti = 4.5, 6, 12, 14.5 at.%) and three had Ti/Si > 1 (range 1.1 to 1.47, Ti = 18, 20, 23.5 at.%). The microstructures of the alloys were studied using EPMA and XRD. One of their alloys, namely Nb-23.5Ti-16Si-10B, was close to the actual composition of the alloy TT1 [12] and its microstructure was reported to consist of Nb_{ss}, T2, Nb₃Si and Nb₃B₂ compared with Nb_{ss}, T2 and Nb₃Si for TT1. It should also be noted that the Nb₃B₂ compound was not reported in the microstructure of the Nb-22Ti-16Si-3Al-5Cr-4Hf-10B (at.%) alloy [29]. According to the EPMA data of Sun et al. [11], in the seven Nb-Ti-Si-B alloys that they studied, the Nb₃B₂ had 39.7 < Si + B < 100042.2 at.% (average 41 at.%), 0.5 < Si/B < 0.76 (average 0.64), 1.9 < Ti < 11.4 at.% (average 6.2 at.%) and 1.37 < (Nb + Ti)/(Si + B) < 1.52 (average 1.44). The T2 silicide had 33.8 < Si + 1.52B < 38.9 at.% (average 37.2 at.%), 1 < Si/B <8.3 (average 3.5), 6.5 < Ti < 18 at.% (average 14 at.%) and 1.6 < (Nb + Ti)/(Si + B) < 2 (average 1.7). Differences for the Si/B ratio and Ti content in Nb₃B₂ and D8₈, and Ti content in T2, for the alloys studied by Sun et al. [11] and this work should be noted. In all four alloys of this work, the T2 was the primary phase. Considering the data for the alloy TT1 [12] and this work, there was two-phase Nb_{ss} and T2 equilibria in TT3, three-phase Nb_{ss}, T2 and Nb₃Si equilibria in TT1 and three-phase Nb_{ss}, T2 and D8₈ equilibria in TT8. Only the synergy of B with Al and Al + Cr in the alloys TT3 and TT4 was able to control the contamination of the microstructure by nitrogen (Table 4). The Si content in the Nb_{ss} increased with its Ti concentration (Figure 4a) and was essentially the same for all the HT alloys, about 0.5 at.%, the latter in agreement with other work on B-free RM(Nb)ICs e.g., [16,17]. Whereas the Si concentration was the same, the B content on the HT alloys varied depending on element(s) in synergy with B (Figure 4b). In the Ti-rich Nb_{ss} and "normal" Nb_{ss} the maximum B concentration respectively was about 6.1 and 3.2 at.%, which corresponded respectively to 1.7 and 1.9 at.%Si. The concentrations of B, Cr and Ti depended on each other, the Ti and B concentrations in the Nb_{ss} respectively increased and decreased with increasing Cr content in the Nb_{ss} (Figure 4c). As the Al + B + Ti content of the Nb_{ss} decreased the VEC of the solid solution increased, whereas the opposite was the case regarding B + Cr + Ti (Figure 4d). The VEC of the Nb_{ss} increases with Ti + Cr or Ti content in the alloys TT2, TT4 and TT8 (Figure 4e,f). The B or Mo content of the Nb_{ss} decreased with increasing Ti concentration in the Nb_{ss} (Table 5).



Figure 4. Cont.



Figure 4. Cont.



Figure 4. (a) Ti versus Si and (b) B versus Si content of the Nb_{ss} in the alloys TT1 [12], TT2, TT3, TT4 and TT8 where the data for HT alloys is shown in green. In (a) the R² values are given for the linear or parabolic fit of data and in (b) the R² values are for the parabolic fit of data, where blue filled circles correspond to Ti-rich Nb_{ss} and red to normal Nb_{ss} in AC alloy. (c) Element X versus Cr in Nb_{ss}, data for X = B (brown) or Ti (purple) in the Nb_{ss} of the HT alloys TT2, TT4 and TT8. (d) Ti + B + X (X = Al, Cr) versus VEC of Nb_{ss} in the HT alloys TT2, TT3, TT4 and TT8, purple X = Al, alloys TT3, TT4, TT8, pink X = Cr, alloys TT2, TT4, TT8, (e) Ti + Cr versus VEC and (f) Ti versus VEC of Nb_{ss}, HT alloys TT2, TT4, TT8.

In the T2 silicide, the B content decreased and the Al and Cr concentrations increased as the Ti concentration increased ([6] and Table 5). Figure 5 shows maps of the T2 silicide in the AC and HT alloys TT2, TT3, TT4 and TT8 and of the Nb₅Si₃ silicide in the reference B-free alloys KZ4, KZ5, KZ7 and JG3. The T2 occupies a distinct, separate area in the $\Delta\chi$ versus VEC map (Figure 5a,b) in agreement with [2,6]. Both VEC and $\Delta\chi$ increase with increasing <Nb> and the linear fit of the data for T2 and Nb₅Si₃ is remarkably good (Figure 5c,d).



Figure 5. Cont.



Figure 5. Maps of the 5-3 silicide in the AC and HT boron-containing alloys TT2, TT3, TT4 and TT8, i.e., of the T2 silicide, and the reference B-free alloys KZ4, KZ5, KZ7 and JG3, i.e., of Nb₅Si₃. (**a**) and (**b**) $\Delta \chi$ versus VEC, (**c**) VEC versus <Nb> and (**d**) $\Delta \chi$ versus <Nb>. Diamonds for B-containing alloys, i.e., data for T2, and pink filed circles for the B free alloys, i.e., data for Nb₅Si₃. (**a**) and (**d**) data for T2, Ti-rich T2, Nb₅Si₃ and Ti-rich Nb₅Si₃. (**b**) and (**c**) data for T2 and Nb₅Si₃ in the aforementioned alloys. In (**b**) and (**c**) green for TT2 and KZ4, red for TT3 and KZ7, purple for TT4 and KZ5, and blue for TT8 and JG3. <Nb> means Nb and other elements that substitute Nb in silicide. For all data in (**a**) R² = 0.5173 for linear fit. The dashed lines in (**a**) delineate the "territory" occupied by the T2 silicide in the B containing alloys TT2, TT3, TT4 and TT8 (defined by $\Delta \chi$ and VEC approximately less than 0.303 and 4.36).

4.3. Hardness

The hardness of the Nb_{ss} decreased or increased with the parameter VEC_{Nbss} when the alloys TT4 and TT8 were grouped respectively with the Cr containing alloy TT2 (Figure 6a) or the Al-containing alloy TT3 (Figure 6b). The same was the case for the reference alloys KZ5 and JG3 when grouped with the Cr or Al-containing alloys KZ4 and KZ7, respectively (Figure 6). The hardness of the Nb_{ss} in the AC and HT alloys TT2, TT4 and TT8 decreased with increasing VEC_{Nbss} (Figure 6c).

The dependence of the hardness of the Nb_{ss} on solutes in the alloys is shown in Figure 7. The hardness decreased with increasing Ti + B + X or Ti + X content (X = Al, Cr), and this was linked with changes of the parameter VEC_{Nbss} (Figures 4 and 6), and also decreased with increasing Ti content in the Nb_{ss}, the same trend as for the reference alloys (Figure 7d). Note the link between Ti in the solid solution and its VEC (Figure 4f) and the change of hardness with VEC (Figure 6a).



Figure 6. Vickers hardness (HV) of Nb_{ss} in HT alloys versus the parameter VEC of the solid solution. (a) alloys TT2, TT4, TT8 (red), KZ4, KZ5 and JG3 (blue), all data $R^2 = 0.7871$ (b) alloys TT3, TT4, TT8 (red), KZ7, KZ5 and JG3 (blue), all data $R^2 = 0.8277$. (c) Vickers hardness of Nb_{ss} in AC (green) and HT (purple) alloys TT2, TT4, TT8, all data $R^2 = 0.8962$.



Figure 7. Cont.



Figure 7. Vickers hardness (HV) of Nbss in the HT alloys TT2, TT3, TT4, TT8 versus (**a**) Ti + B + X, (**b**) Ti + X, (X = Al (blue), Cr (red)), (**c**) Ti/Cr for the alloys TT2, TT4, TT8, and (**d**) HV of Nbss in the HT alloys TT3, TT4, TT8 (green) and KZ7, KZ5 and JG3 (brown).

The VEC_{T2} or $\Delta\chi_{T2}$ increased with increasing <Nb> in T2 (Figure 5c,d) and the hardness of T2 increased with VEC_{T2} (in [2]). The synergy of Cr with B and Ti decreased slightly the hardness of T2 in TT2-AC, which was the highest amongst the AC alloys of this study, compared with TT1-AC, but the T2 in TT2-HT did not retain its hardness after heat treatment, which decreased by 13% compared with 6% in TT1-HT ([12] and Table 2) and was essentially the same as the hardness of T2 in TT3-HT and the second-lowest amongst all the HT alloys (Table 2). Best hardness retention of the T2 and highest hardness after heat treatment was observed in the case of the alloy TT4. Compared with the alloys ZF4 [30], ZF5 [31] and ZF6 [32], which are similar respectively to TT2, TT3 and TT4 with the addition of Ge instead of B, the T2 had significantly lower hardness than Nb₅Si₃ in these three alloys. Compared with TT1, the T2 hardness in TT4-HT was not significantly different than that in TT1-HT [12], which was also lower than the Nb₅Si₃ in ZF3 [33]. It should be noted that the relationship between the hardness of T2 and VEC_{T2} is opposite that of Nb₅Si₃ and VEC_{Nb5Si3} (in [2]).

Alloy hardness versus the parameters VEC_{alloy}, δ_{alloy} and $\Delta \chi_{alloy}$ is shown in Figure 8a–c. Room temperature specific strength calculated from hardness $(\sigma_{HV}/\rho)_{alloy}$ is shown in Figure 8d. The addition of B to the reference alloys caused a decrease in VEC_{alloy} and an increase in HV_{alloy} and $(\sigma_{HV}/\rho)_{alloy}$. Notice that the B containing alloys occupy specific distinct areas in the HV_{alloy} versus δ_{alloy} or $\Delta \chi_{alloy}$ maps.



Figure 8. Cont.



Figure 8. (a) to (c) Vickers hardness (HV) and (d) room temperature specific strength calculated from hardness (σ_{HV}/ρ) versus the parameters (a), (d) VEC, (b) δ and (c) $\Delta\chi$ of the TT2, TT3, TT4, TT8, KZ4, KZ5, KZ7 and JG3 alloys. In (a) to (c) the data is for the AC and HT conditions and in (d) the data is for the AC condition. Filled circles green TT2 and KZ4, red TT3 and KZ7, purple TT4 and KZ5, and blue TT8 and JG3. In (b) and (c) ellipses enclose the B containing alloys. In (a) and (d) the R² values are for the parabolic fit of data. In (a) data for the alloys TT2, TT3, TT4 and TT8 with R² = 0.998 and R² = 0.9046 for the AC and HT conditions, respectively and with R² = 0.9922 and R² = 0.9908 respectively for the AC and HT alloys KZ4, KZ5, KZ7 and JG3. In (d) R² = 0.9993 and R² = 0.9888 respective for the AC alloys TT2, TT3, TT4, TT8 and KZ4, KZ5, KZ7, JG3. In (a) and (d) green arrows indicate the effect of B addition.

Furthermore, notice the location of the JG3 alloy in the HV_{alloy} versus $\Delta \chi_{alloy}$ map. The maximum specific strength of 385 MPa cm³g⁻¹ for VEC_{alloy} = 4.45 from the parabola for B containing alloys in Figure 8d is significantly higher than those reported for RCCAs to date [1,2], and only slightly lower than that of the RM(Nb)ICs-RCCAs JZ4 and JZ5 [34].

The parameters VEC_{alloy}, δ_{alloy} and $\Delta \chi_{alloy}$ link with specific solutes and thus allow us (i) to link the hardness and specific strength of alloys with solute additions and; (ii) to use these relationships for the design of alloys using the alloy design methodology NICE [1–3]. We demonstrate such relationships for the alloys TT2, TT4 and TT8 and the solutes B, Cr and Ti in Figure 9.



Figure 9. Cont.



Figure 9. Cont.



Figure 9. Cont.



Figure 9. Maps of the parameters VEC_{alloy}, δ_{alloy} and $\Delta \chi_{alloy}$ with sums or ratios of Ti, Cr or B in AC and HT alloys TT2, TT4, TT8 and "equivalent" B free alloys KZ4, KZ5 and JG3. R² values for linear or parabolic fit of data. (a) VEC_{alloy} versus (Ti + Cr)_{alloy}, (b) δ_{alloy} versus (Ti + Cr)_{alloy}, (c) $\Delta \chi_{alloy}$ versus (Ti + Cr)_{alloy}, (d) VEC versus Ti/Cr, (e) δ versus Ti/Cr, (f) $\Delta \chi$ versus Ti/Cr, (g) VEC versus Ti + Cr + B, (h) δ versus Ti + Cr + B and ((i) $\Delta \chi$ versus Ti + Cr + B. (a) to (f) data for the alloys TT2, KZ4, TT4, KZ5, TT8 and JG3, (g) to (i) data for the alloys TT2, TT4, TT8. (a,b,d,e) colours as follows: green for TT2 and KZ4, purple for TT4 and KZ5, and blue for TT8 and JG3. (g) to (i) triangles for TT8, squares for TT4, and circles for TT2. (c) and (f) green colour for all alloys. In (a) to (f) diamonds for B containing alloys.

The parameter VEC_{alloy} increased with (Ti + Cr)_{alloy} (Figure 9a) whereas the parameters δ_{alloy} and $\Delta \chi_{alloy}$ decreased (Figure 9b,c). The shift towards higher HV_{alloy} with decreased VEC_{alloy} for the B-containing alloys (Figure 8a) was thus attributed to the lower (Ti + Cr) content in the alloys TT2, TT4 and TT8. The shift of the latter in separate distinct areas in the maps in Figure 8b,c was attributed to the increase of δ_{alloy} and $\Delta \chi_{alloy}$ with the decrease of (Ti + Cr)_{alloy} (Figure 9b,c). The parameter VEC_{alloy} decreased and the δ_{alloy} and $\Delta \chi_{alloy}$ increased with increasing (Ti/Cr)_{alloy} (Figure 9d,e,f). The alloys were separated in the VEC_{alloy} or δ_{alloy} versus (Ti / Cr)_{alloy} maps, as was the case in the VEC_{alloy} or δ_{alloy} such the former two maps the case for Mo-containing alloys JG3 and TT8 was different, with VEC_{alloy} minimum at about 4.45 and δ_{alloy} maximum at about 15 for (Ti/Cr)_{alloy} = 4.75. Note that VEC_{alloy} ≈ 4.45 corresponds to maximum HV_{alloy} and

specific strength in Figure 8a,d. Similarly to the $\Delta \chi_{alloy}$ versus (Ti + Cr)_{alloy} map, the alloys were not separated in the $\Delta \chi_{alloy}$ versus $(Ti/Cr)_{alloy}$ map (Figure 9c,f). Thus, even though there are relationships of the parameters VEC_{allov}, δ_{allov} and $\Delta \chi_{allov}$ with (Ti + Cr)_{allov}, or $(Ti/Cr)_{alloy}$, and the parameters VEC_{alloy}, δ_{alloy} and $\Delta \chi_{alloy}$ are used to design new alloys using NICE [1–3], only the parameter VEC_{alloy} links hardness or specific strength with specific solutes in NICE, (Ti + Cr)_{alloy} and (Ti/Cr)_{alloy} in the case demonstrated above. Furthermore, the parameters VEC_{alloy}, δ_{alloy} and $\Delta \chi_{alloy}$ also link with the (Ti + Cr + B) content of the alloys TT2, TT4 and TT8, as shown in the maps in Figure 9g,h,i. VECalloy is maximum for $(Ti + Cr + B)_{allov} = 35.5 at.\%$. The HV_{allov} versus $(Ti + Cr + B)_{allov}$ map (not shown) shows minimum (685 HV_{allov}) and maximum (740 HV_{allov}) hardness, respectively, for AC and HT alloys for Ti + Cr + B = 35.5 at.% whereas the HV_{alloy} versus (Ti + Al + B)_{alloy} map (not shown) shows minima of 615 HV_{alloy} and 560 HV_{alloy} with (Ti + Al + B)_{alloy} equal to 34.5 and 35.5 at.%, respectively, for the AC and HT alloys TT3, TT4 and TT8. It should be noted that also the hardness of the solid solution (HV_{Nbss}) links with VEC_{Nbss} (Figure 6) and with solutes, namely Ti + X, Ti + B + X (X = Al, Cr), and Ti/Cr (Figure 7) and that solutes in the Nb_{ss} link with its parameters (see Figure 4 for relationships with VEC_{Nbss} for B containing alloys and Figure A1 in the Appendix A for relationships in the reference alloys). Such relationships allow the design methodology NICE to optimise alloy composition for the balance of strength and oxidation properties, see [1–3].

Figure 10 compares the density of the base alloys KZ4, KZ7 and KZ5 with that of the equivalent alloys with Ge addition, namely ZF4, ZF5 and ZF6 [30–32], and with B addition.



Figure 10. Densities of as cast alloys KZ4, KZ7, KZ5 (dark blue, 1, 4, 7), ZF4, ZF5, ZF6 (red, 2, 5, 8) and TT2, TT3, TT4 (green, 3, 6, 9). See Appendix A for nominal compositions of alloys.

It is clear that the B addition had the most significant effect on reducing alloy density. Figure 11 compares the room temperature hardness of AC and HT alloys ZF4, ZF5, ZF6 [30–32] and TT2, TT3 and TT4. The addition of Ge to the reference alloys increased the alloy hardness compared with the addition of B. This was attributed (i) to the increase of the hardness of the Nb₅Si₃ when alloyed with Ge; and (ii) to the lower vol.% of Nb_{ss} in the Ge containing alloys [6,30–32] compared with the alloys TT2, TT3 and TT4.



Figure 11. Room temperature Vickers hardness of AC and HT alloys. Odd numbers AC alloys, even numbers HT alloys. TT2 (1, 2), ZF4 (3, 4), TT3 (5, 6), ZF5 (7, 8), TT4 (9, 10), ZF6 (11, 12). See Appendix A for nominal compositions of alloys.

4.4. Oxidation

The data for the alloys TT1, TT2, TT3, TT4 and TT8 can help us to infer what the effect of B addition to the reference alloys was on isothermal oxidation at 800 and 1200 °C and to compare the effect of B addition with that of Ge or Sn to the same reference alloys. Amongst the reference alloys KZ4, KZ5, KZ7 and JG3, the alloy JG3-AC had the best oxidation behaviour at both temperatures, did not pest at 800 °C, and followed parabolic oxidation at 800 °C and liner oxidation at 1200 °C (the JG3-HT followed linear oxidation at both temperatures) [35]. The rate constant of the alloys TT1, TT2 and TT3 at 800 °C (Table 6) was one order of magnitude higher than that of JG3-HT, and of the alloys TT4 and TT8 similar or one order of magnitude lower, respectively. The alloys KZ4, KZ5 and KZ7 formed Maltese crosses at 800 °C [36], like the MASC alloy, and their scales spalled off at 1200 °C, as did the scale of JG3.

At 800 °C, the mass change of TT1 was significantly lower than that of KZ3, which gained more than about 200 mg/cm² after just 10 h. The synergy of B and Cr decreased the mass change compared with KZ4 (about 60 mg/cm² after 100 h). The Al addition in TT3 also improved the oxidation behaviour compared with TT1 (Table 6). Compared with KZ7, the synergy of Al with B was only marginally beneficial for the first 20 h, after this time the mass change increased. The mass of the alloy TT4 did not change for the first 10 h and after this time it was lower than that of KZ5, which gained about 32.5 mg/cm². The Mo addition in TT8 significantly improved the oxidation behaviour compared with JG3, which gained about 3.5 mg/cm².

At 1200 °C, the synergy of Al and Cr in TT4 had a strong effect on oxidation behaviour (mass change after 90 h: TT1 = 80.3, TT2 = 105.6, TT3 = 66.1 and TT4 = 34.0 mg/cm²). The addition of Mo in the TT8 alloy resulted in a marginal improvement of the oxidation behaviour of TT8 compared with TT4. Figure 12 compares the mass changes of the alloys of this study with their reference alloys after 90 h at 800 or 1200 °C. In each part of this figure, the effect of B addition is indicated by an arrow. With the exception of TT3, whose mass increased more than KZ7 at 800 °C (Figure 12b), the addition of B resulted to lower mass changes at both temperatures, particularly in the case of the alloy TT8.



Figure 12. Cont.



Figure 12. Comparison of mass changes of B-containing alloys TT2, TT3, TT4 and TT8 with reference alloys KZ4, KZ5, KZ7 and JG3. For isothermal oxidation at 800 and 1200 °C. Arrows indicate direction of mass change at 800 °C (**a**–**d**) and 1200 °C (**e**–**h**).

Plots of the mass changes of the B-containing alloys and their reference alloys versus the alloy parameters VEC and δ are shown in Figure 13. According to the alloy design methodology NICE, for improved oxidation behaviour, the alloy design should aim to lower VEC_{alloy} and increase δ_{alloy} [1–3]. Figure 13 shows that this was the case for all alloys at 1200 °C, and also at 800 °C where the alloy TT3 was the exception as its mass change was higher than that of KZ7 (Figures 12b and 13a,b).



Figure 13. Cont.



Figure 13. Mass change (Δ W/A) of the alloys TT2, TT3, TT4, TT8 and their basis/reference alloys KZ4, KZ5, KZ7 and JG3 versus the parameters (**a**,**c**) VEC and (**b**,**d**) δ for isothermal oxidation (**a**,**b**) at 800 °C and (**c**,**d**) at 1200 °C. Colours as follows: green for TT2 and KZ4, red for TT3 and KZ7, purple for TT4 and KZ5, and blue for TT8 and JG3. Diamonds for B containing alloys.

Compared with the Ge containing alloys ZF4, ZF5 and ZF6 [37] and the Sn containing alloys ZX4, ZX6 and ZX8 [22] (for nominal compositions see Table A1 in the Appendix A), the oxidation of the B-containing alloys TT2, TT3 and TT4 was inferior at 800 °C. Indeed, the above ZF and ZX series alloys did not pest, their scales did not spall off, and followed parabolic oxidation kinetics [37], or parabolic with/without linear kinetics [22]. At 1200 °C the ZF and ZX series alloys followed either parabolic plus linear kinetics [37] or parabolic or linear kinetics [22] and their scales spalled off, the same as the KZ series reference alloys. Only the alloy TT8 did not pest at 800 °C and its scale did not spall off at 800 and 1200 °C.

At 800 °C, the scales that formed on the alloys TT1 and TT2 consisted of SiO₂, TiO₂, Nb₂O₅, 3Nb₂O₅.TiO₂, and 5Nb₂O₅.2TiO₂, which are typical oxides in the scales of RM(Nb)ICs [3,35,36], plus B₂O₃ and B₂SiO₅ in TT1 plus CrNbO₄ in TT2 (Figure S2a in Supplemental data). The scales that formed on TT3 and TT4 consisted of the same oxides as TT1, plus AlNbO₄ in TT3 and CrNbO₄ and AlNbO₄ in TT4 (data not shown). At 1200 °C, the scale on TT1 consisted of SiO₂, TiO₂, Nb₂O₅ and 3Nb₂O₅.TiO₂, 5Nb₂O₅.2TiO₂, plus B₂O₃ and B₂SiO₅ whereas the scale on TT2 consisted of the same oxides plus NbO, and CrNbO₄ (Figure S2b in Supplementary Materials). The scale of TT3 consisted of the same oxides as TT1 plus AlNbO₄ (data not shown). The presence of CrNbO₄ and AlNbO₄ in the scales of the alloys TT2, TT3 or TT4 is in agreement with the data for the B free alloys KZ4, KZ7 and KZ5 [3,36] and the Ge containing alloys ZF4, ZF5 and ZF6, where also SiO₂, Nb₂O₅, TiNb₂O₇ (Nb₂O₅.TiO₂) and GeO₂ were observed [37].

Comparison of the oxidation of the reference alloys KZ4, KZ7, KZ5 [36] with the alloys with B addition, namely TT2, TT3 and TT4, or Ge addition (ZF4, ZF5, ZF6 [37]) or Sn addition (ZX4, ZX6, ZX8 [22]) shows that at 800 °C (i.e., in pest regime) Ge and B, respectively, was most and least effective and that at 1200 °C all three elements were not effective as scale spallation was not suppressed. However, only when B was in synergy with Al, Cr and Mo was the oxidation improved compared with the basis alloy JG3 [35], as the alloy TT8 did not pest and its scale did not spall off at 800 and 1200 °C. Thus, the aforementioned synergy of elements in TT8 had the same effect at both temperatures as that of Al and Cr with Ge plus Sn in the alloy OHS1 [23]. In other words, the alloy TT8 had outstanding (for RMICs, and RHEAs and RCCAs) oxidation behaviour at 800 and 1200 °C with density 6.4 g/cm³ and specific strength of 377 and 337 MPacm³g⁻¹, respectively, in AC and HT conditions. Another characteristic difference of the alloys with B or Ge addition [37] with the Sn containing alloys ZX4, ZX6 and ZX8 [22] was the fact that in the latter, tin oxide was rarely (if at all) observed in the scale, in contrast with the former alloys where B₂O₃ or GeO₂ were formed. The latter two oxides form glass with silica.

5. Conclusions

We studied the effect of B addition on the density, macrosegregation, microstructure, hardness and oxidation of four RM(Nb)IC alloys, namely the alloys TT2, TT3, TT4 and TT8. In actual fact, these alloys were based on four KZ series alloys (the basis/reference alloys) to which B was added. This choice of alloys made it possible to also compare the effect of B addition on density, hardness or oxidation with that of Ge or Sn addition.

The addition of B resulted in B macrosegregation that was highest in TT8 and in increased macrosegregation of Si and Ti, both of which were lowest in TT8. With the B addition, the density of the alloys was decreased significantly and was lower than the alloys with Ge addition. The alloy TT8 had the lowest density of 6.41 g/cm³ and the highest specific strength at room temperature, which was also higher than that of RCCAs and RHEAs. The Nb_{ss} and T2 silicide were stable in the alloys TT2 and TT3, whereas in TT4 and TT8 the stable phases were the Nb_{ss} and the T2 and D8₈ silicides. The latter silicide was Al free. The T2 had Si + B + Al \approx 37.5 at.%, Si/B \approx 4 and <Nb>/<Si> \approx 1.7 compared with Si + B \approx 41.5 at.%, Si/B \approx 0.5 and <Nb>/<Si> \approx 1.4 of the D8₈. Compared with the Ge or Sn addition in the same reference alloy, the B and Ge addition was the least and most effective at 800 °C (i.e., in the pest regime), when no other RM was present in the

alloy. Like Ge or Sn, the B addition in TT2, TT3 and TT4 did not suppress scale spallation at 1200 °C. Only the alloy TT8 did not pest and its scales did not spall off at 800 and 1200 °C.

The macrosegregation of Si and Ti, the chemical composition of Nb_{ss} and T2, the microhardness of Nb_{ss} and the hardness of alloys, and the oxidation of the alloys at 800 and 1200 °C were also viewed from the perspective of the alloy design methodology NICE and relationships with the alloy or phase parameters VEC, δ and $\Delta \chi$. The trends of these parameters and the location of alloys and phases in parameter maps were found to be in agreement with NICE, the one exception being the macrosegregation of Si in TT8 when compared with its basis/reference alloy.

Supplementary Materials: The following are available online at https://www.mdpi.com/article/ 10.3390/ma14206101/s1, Figure S1: X-ray diffractograms (a), (c), (e) and (g) of AC alloys TT2, TT3, TT4 and TT8, respectively, and (b), (d), (e) and (h) of HT alloys TT2, TT3, TT4 and TT8, respectively; Figure S2: X-ray diffractograms of oxide scale formed on the alloy TT2 (a) at 800 °C and (b) at 1200 °C.

Author Contributions: Experimental work, T.T.; funding, P.T.; supervision, P.T.; formal analysis, T.T. and P.T.; draft preparation, T.T.; review, P.T., final paper, P.T. All authors have read and agreed to the published version of the manuscript.

Funding: This research was funded by the EPSC (EP/H500405/1, EP/L026678/1).

Institutional Review Board Statement: Not applicable.

Informed Consent Statement: Not applicable.

Data Availability Statement: All the data for this paper is given in the paper and its Supplementary Materials, other data cannot be made available to the public.

Acknowledgments: The support of this work by the University of Sheffield, Rolls-Royce Plc and EPSRC (EP/H500405/1, EP/L026678/1) is gratefully acknowledged.

Conflicts of Interest: The authors declare no conflict of interest.

Appendix A

$$\delta = 100 \sqrt{\sum_{n=1}^{n} c_{\rm i} \left(1 - r_{\rm i}/\bar{r}\right)^2}$$
 (A1)

$$\Delta \chi = \sqrt{\sum_{n=1}^{n} c_{\rm i} \left(\chi_{\rm i} - \overline{\chi}\right)^2} \tag{A2}$$

$$VEC = \sum_{n=1}^{n} c_{i} (VEC)_{i}$$
(A3)

$$\bar{r} = \sum_{n=1}^{n} c_{i} r_{i}$$
(A4)

$$\overline{\chi} = \sum_{n=1}^{n} c_{i} \chi_{i}$$
(A5)

$$\Delta H_{mix} = \sum_{i=1, j \neq 1}^{n} \prod_{ij} c_i c_j \tag{A6}$$

$$\Pi_{ij} = 4\Delta_{mix}{}^{AB} \tag{A7}$$

$$\Delta S_{mix} = -R \sum_{i=1}^{n} c_i \ln c_i \tag{A8}$$

$$\mathbf{T}_m = \sum_{i=1}^n c_i \, \mathbf{T}_{\mathbf{m}_i} \tag{A9}$$

where c_i , r_i , χ_i , (*VEC*)_{*i*} and T_{mi} , respectively, are atomic percentage, atomic radius, Pauling electronegativity, VEC and melting point of *i*th element, Δ_{mix}^{AB} is the mixing enthalpy of binary liquid *AB* alloy and *R* is the gas constant.

A 11							Element	:		
Alloy	Nb	Ti	Si	Al	Cr	Мо	В	Ge	Sn	Ref.
JG3	bal.	24	18	5	5	2	-	-	-	[17]
KZ3	bal.	24	18	-	-	-	-	-	-	[16]
KZ4	bal.	24	18	-	5	-	-	-	-	[16]
KZ5	bal.	24	18	5	5	-	-	-	-	[16]
KZ7	bal.	24	18	5	-	-	-	-	-	[16]
OHS1	bal.	24	18	5	5	-	-	5	5	[23]
TT1	bal.	24	18	-	-	-	8	-	-	[12]
ZF3	bal.	24	18	-	-	-	-	5	-	[33]
ZF4	bal.	24	18	-	5	-	-	5	-	[30]
ZF5	bal.	24	18	5	-	-	-	5	-	[31]
ZF6	bal.	24	18	5	5	-	-	5	-	[32]
ZX4	bal.	24	18	-	5	-	-	-	5	[22]
ZX6	bal.	24	18	5	-	-	-	-	5	[22]
ZX8	bal.	24	18	5	5	-	-	-	5	[22]

Table A1. Nominal compositions (at.%) of reference alloys used in this work.

Table A2. $\langle Nb \rangle / \langle Si \rangle$ and Si / B ratios and Si + B and Si + B + Al sums in T2 and D88 silicides in AC and HT alloys, where $\langle Nb \rangle = Nb + TM + RM$, $\langle Si \rangle = Al + B + Si$, TM = Cr, Ti and RM = Mo.

Allow and Condition	Dhaaa	Parameter						
Alloy and Condition	Phase –	Si + B (at.%)	Si/B	Si + B + Al (at.%)	<nb <si=""></nb>			
TT2-AC	T2	38.2	2	-	1.6			
-	Ti rich T2	37.1	3.8	-	1.7			
TT2-HT	T2	37.2	1.9	-	1.7			
-	Ti rich T2 *	34.1 to 37 Average 35.6	1, 3.8, 6.3 Average 3.7	-	1.7 to 1.9 Average 1.8			
TT3-AC	T2	-	1.9	38.6	1.6			
-	Ti rich T2	-	4	39	1.56			
-	T2 very rich in Ti	-	5	37.5	1.67			
TT3-HT	T2	-	2.6	38.5	1.6			
-	Ti rich T2	-	6.1	37.7	1.65			
TT4-AC	T2	-	4.3	38.2	1.62			
-	Ti rich T2	-	6.4	36.7	1.72			
-	D88	40.8	0.6	-	1.45			
TT4-HT	T2	-	5.5	39	1.56			
-	D8 ₈	42.7	0.5	-	1.3			
TT8-AC	T2	-	2.5	38.9	1.57			
-	Ti rich T2	-	3.3	37.7	1.65			
-	D8 ₈	42.6	0.4	-	1.35			
TT8-HT	T2	-	6.2	37.3	1.68			
-	Ti rich T2	-	4.1	38.3	1.61			
-	D88	40	0.5	-	1.5			



Figure A1. Data for the Nb_{ss} in HT boron free alloys KZ4, KZ5, KZ7 and JG3. (**a**)Ti + X (X = Al (blue), Cr (red)) versus the parameter $\Delta \chi$ for the alloys KZ4, KZ5, KZ7 and JG3, (**b**) and (**c**) Ti/Cr versus (**b**) δ for the alloys KZ4, KZ5 and JG3 and (**c**) $\Delta \chi$ for the alloys KZ7, KZ5 and JG3.

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