

MAGNETIC PROPERTIES OF TETRATAENITE-RICH METEORITES II

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Abstract: Magnetic hysteresis and thermomagnetic characteristics of St. Séverin (LL6), Appley Bridge (LL6) and Tuxtuac (LL5) chondrites, which contain tetraetaenite in their metallic components, are measured and analyzed in comparison with another tetraetaenite-rich chondrite, Yamato-74160 (LL7). The magnetic properties of tetraetaenite-rich meteorites are characterized by (a) high magnetic coercive force (H_c) which amounts to 520 Oe for St. Séverin and 160 Oe for Appley Bridge, (b) essential flatness up to about 500°C and then a sharp irreversible drop down to Curie point of the first-run heating thermomagnetic curve. Both characteristic features are broken down to the ordinary features of disordered taenite by a breakdown of tetraetaenite structure at elevated temperatures beyond the order-disorder transition temperature.

The natural remanent magnetization (NRM) of tetraetaenite-rich meteorites is extremely stable against AF-demagnetization and other magnetic disturbances because of the high magnetic coercivity of tetraetaenite. The breakdown processes of ordered tetraetaenite structure by heat treatments are experimentally pursued to understand the formation process of tetraetaenite phase in meteorites.

1. Introduction

In previous papers (NAGATA and FUNAKI, 1982, 1983), magnetic properties of Yamato-74160 LL7 chondrite, Allan Hills-77260 L3 chondrite, St. Séverin LL6 chondrite and ALH-77219 mesosiderite, which contain the tetraetaenite phase in their metallic components, have been reported. As the magnetic coercivity of tetraetaenite is extremely high because of its large magnetic anisotropy owing to its tetragonal crystal structure, the natural remanent magnetization (NRM) of tetraetaenite-rich meteorites is exclusively stable against AF-demagnetizations or similar disturbances. It seems likely therefore that tetraetaenite-rich meteorites are uniquely important in paleomagnetic studies of the primordial solar system.

CLARKE and SCOTT (1980) pointed out that they have optically detected the tetraetaenite phase in 15 H chondrites, 8 L chondrites, 9 LL chondrites, 18 mesosiderites, 2 pallasites, 6 irons and a diogenite. Among these meteorites which are optically proved to contain the tetraetaenite phase, ALH-77219 mesosiderite (NAGATA and FUNAKI, 1983), Estherville mesosiderite and Appley Bridge LL6 chondrite (WASILEWSKI, 1982) are examined magnetically also to detect presence of tetraetaenite in them. The tetraetaenite phase in Y-74160 and ALH-77260 chondrites has been detected on the basis of its optical anisotropy in addition to its characteristic chemical composition of

50%Fe50%Ni in atomic ratio (NAGATA and FUNAKI, 1982, 1983), while that in St. Séverin chondrite on the basis of Mössbauer spectra (DANON *et al.*, 1979a, b), before characteristic behaviours of magnetic properties caused by the presence of tetrataenite phase in these meteorites are magnetically examined.

The characteristic magnetic behaviours of tetrataenite phase observed so far in these tetrataenite-rich meteorites may be summarized by two main features, namely (a) a constancy of saturation magnetization intensity with a very small decreasing rate with increasing temperature up to about 450°C, which is irreversibly broken down by heating above Curie point, and (b) an unusually high magnetic coercivity which also is broken down to a low coercivity state of the disordered taenite caused by a breakdown of the anisotropic ordered structure by heating above Curie point. It appears, however, that the magnetic behaviour of tetrataenite phase in meteorites is fairly complicated owing to coexistence of disordered taenites and kamacites of various Ni contents, which are in close contact with the tetrataenite phase in individual metallic grains (*e.g.* NAGATA and FUNAKI, 1982).

Since the formation and breakdown processes of tetrataenite phase and necessary conditions for its equilibrium state in meteorites are not fully clarified yet, descriptions in detail of mineralogical and magnetic properties of as many tetrataenite-rich meteorites as available will be still required in the next step of studies. In the present work, magnetic properties of St. Séverin (LL6), Appley Bridge (LL6) and Tuxtuac (LL5) tetrataenite-rich chondrites will be examined with reference to their metallic compositions which are estimated by Mössbauer spectral analyses and other methods. In particular, St. Séverin chondrite is magnetically examined in some detail, because only metallic grains of St. Séverin were magnetically examined in the previous work. In the present report, bulk specimen, matrix, clast, metallic grains and apparently pure tetrataenite phase specimens extracted from whole metallic grains of St. Séverin chondrite are separately examined.

2. St. Séverin LL6 Chondrite

2.1. Composition and structure of metallic component

Results of Mössbauer spectral analysis of metallic component and an almost pure tetrataenite phase extracted from the metal of St. Séverin are represented by relative abundances of tetrataenite (TT), disordered taenite of less than 30% in Ni content which is paramagnetic at room temperature (PT) and kamacite (K) as

$$(TT):(PT):(K)=51:9.5:39.5 \quad (\text{in wt}\%)$$

for the whole metal, and

$$(TT):(PT):(K)=74:26:0 \quad (\text{in wt}\%)$$

for almost pure tetrataenite phase which is extracted by melting kamacite phases by diluted HCl.

Chemical compositions of opaque minerals in St. Séverin are newly measured with the aid of an electron microprobe analyzer, results being summarized in Table 1, where chemical compositions of taenite, kamacite and troilite are separately summarized. As shown in Table 1, the average composition of taenite phase is given by $(55.92 \pm 2.4)\text{wt}\%$ Fe, $(44.39 \pm 1.8)\text{wt}\%$ Ni and $(0.78 \pm 0.31)\text{wt}\%$ Co, which are con-

Table 1. Chemical composition of opaque minerals in St. Séverin.

Phase	Fe	Ni (wt %)	Co	S	Total
Taenite	59.24	41.30	0.79	0.00	101.33
"	57.37	43.24	0.61	0.00	101.21
"	53.33	45.09	0.29	0.00	98.71
"	52.56	45.45	0.51	0.03	98.55
"	58.25	40.29	1.38	0.00	99.92
"	54.80	44.95	1.08	0.00	100.83
Kamacite	93.36	5.25	2.80	0.01	101.42
"	91.85	5.13	4.20	0.01	101.19
"	91.23	5.15	3.54	0.04	99.96
"	89.76	5.12	3.74	0.01	98.63
Troilite	65.80	0.09	0.06	33.15	99.10
"	62.19	0.09	0.63	36.40	99.31

siderably deviated from the chemical composition of stoichiometric tetrataenite, *i.e.* 48.76wt%Fe, 51.24 wt%Ni. DANON *et al.* (1979a) have already reported that grain sizes of the ordered tetrataenite phase in paramagnetic matrix of Ni-poor taenite in St. Séverin metallic components are ranged mostly from 0.15 to 0.3 μm . It seems likely therefore that the chemical compositions of taenite phase obtained by the electron microprobe analyzer of 3 μm in beam diameter represent, in most cases, average values of chemical compositions of the fine tetrataenite grains and the surrounding Ni-poor taenite. Actually, scanning surveys of low accuracy of the chemical composition of taenite surface of St. Séverin metallic grains have occasionally shown a group of composition such as (50.0%Fe, 52.3%Ni, 0.4%Co) or (45.9%Fe, 53.1%Ni, 3.1%Co) and another group of composition such as (63.3%Fe, 33.9%Ni, 0.0%Co), or (62.6%Fe, 34.3%Ni, 1.2%Co). The former group may represent a tetrataenite-rich spot, while the latter group a low-Ni taenite matrix spot.

The average chemical composition of kamacite phase in St. Séverin is given by $(91.55 \pm 1.06)\text{wt}\%\text{Fe}$, $(5.16 \pm 0.04)\text{wt}\%\text{Ni}$, $(3.57 \pm 0.40)\text{wt}\%\text{Co}$. The chemical composition of kamacite phase is much more homogeneous than that of taenite phase, as expected.

2.2. Basic magnetic properties

Basic magnetic properties which characterize ferromagnetic hysteresis curves are measured at room temperature and at liquid He temperature for the bulk specimen, matrix, clast, metallic grains and apparent tetrataenite phase extracted from the metals of St. Séverin. The magnetic measurements are carried out before the meteorite samples are heated up to elevated temperatures beyond their Curie point and after the heating procedure. Saturation magnetization (I_s), saturated isothermal remanent magnetization (I_R), coercive force (H_C) and remanence coercive force (H_{RC}) are the measured basic ferromagnetic parameters of these samples, as summarized in Table 2.

It is noted first in Table 2 that both H_C and I_R of all samples of St. Séverin are much reduced after heating in comparison with their initial values before the heating,

Table 2. Basic magnetic properties of bulk, matrix, clast, metallic grains and extracted taenite of St. Séverin.

Sample	I_s (emu/g)		I_R (emu/g)		H_C (Oe)		H_{RC} (Oe)	
	Before Heating	After Heating	Before Heating	After Heating	Before Heating	After Heating	Before Heating	After Heating
Bulk								
300 K	2.80	3.95	0.50	0.021	520	9.5	1840	110
4.2 K	3.85	5.10	0.68	0.125	430	39	2680	780
Matrix								
300 K	7.4	10.2	1.43	0.17	590	35	1945	210
4.2 K	8.2	11.7	1.64	0.21	575	38	2240	380
Clast								
300 K	3.21	5.55	0.40	0.14	340	38	1310	300
4.2 K	6.55	6.35	0.60	0.17	335	48	2110	350
Metal								
300 K	142	124	25	0.20	860	3.5	895	47
4.2 K	135	—	24	—	895	—	2300	—
Tetrataenite								
300 K	127	128	23.5	0.40	765	3.5	1480	60
4.2 K	—	144	—	0.70	—	7.0	—	38

indicating that the original ordered tetrataenite structure is broken down to the disordered taenite structure by the heating procedure. It will be further noted that ratios

$$\{I_R(\text{Before heating})/I_R(\text{After heating})\},$$

$$\{H_C(\text{Before heating})/H_C(\text{After heating})\}$$

and

$$\{H_{RC}(\text{Before heating})/H_{RC}(\text{After heating})\}$$

observed at room temperature are particularly large for metals and extracted tetrataenite phase metals in comparison with those for bulk, matrix and clast samples. This result may suggest that the breakdown phenomenon of ordered tetrataenite structure in bulk, matrix and clast samples by heating is associated with some chemical and structural reactions between tetrataenite phase and non-tetrataenite components at elevated temperatures, resulting thus in relaxing the sharp breakdown phenomenon of the magnetically coercive tetrataenite structure at elevated temperatures. As already discussed (NAGATA and FUNAKI, 1982), magnetic coercive force (H_C) of tetrataenite crystal itself along its easy magnetization axis is theoretically estimated to be about 4.9×10^3 Oe and the observed values of H_C smaller than 10^3 Oe of all samples shown in Table 2 are due to coexistence of ferromagnetic component(s) of much lower coercivity.

It may be remarked that matrix and clast in Table 2 are not those which are extracted from the same bulk sample given in the same table. Since the distribution of metallic grains in St. Séverin is fairly heterogeneous, the I_s values of its bulk, matrix and clast specimens can be different from one another by factor 2. For example, the ferromagnetic hysteresis parameters at room temperature of a bulk sample of St. Séverin examined in a preliminary study (NAGATA and FUKUI, 1982) are given by $I_s=4.07$ emu/g, $I_R=0.54$ emu/g, $H_C=500$ Oe and $H_{RC}=1900$ Oe. In regard to Table 2, therefore, ferromagnetic parameters related to magnetic coercivity such as I_R/I_s , H_C and

H_{RC} will be the main interest in the present study.

2.3. Thermomagnetic curve characteristics

The first-run thermomagnetic curves of a bulk sample of St. Séverin in a temperature range between -269 and 850°C in a magnetic field of 10 kOe are shown in Fig. 1. The cooling thermomagnetic curve of the second-run is approximately same as the first-run cooling thermomagnetic curve, though the second-run heating thermomagnetic curve also has a kamacite magnetization component which is represented by $\alpha \rightarrow \gamma$ transition point of kamacite at 780°C .

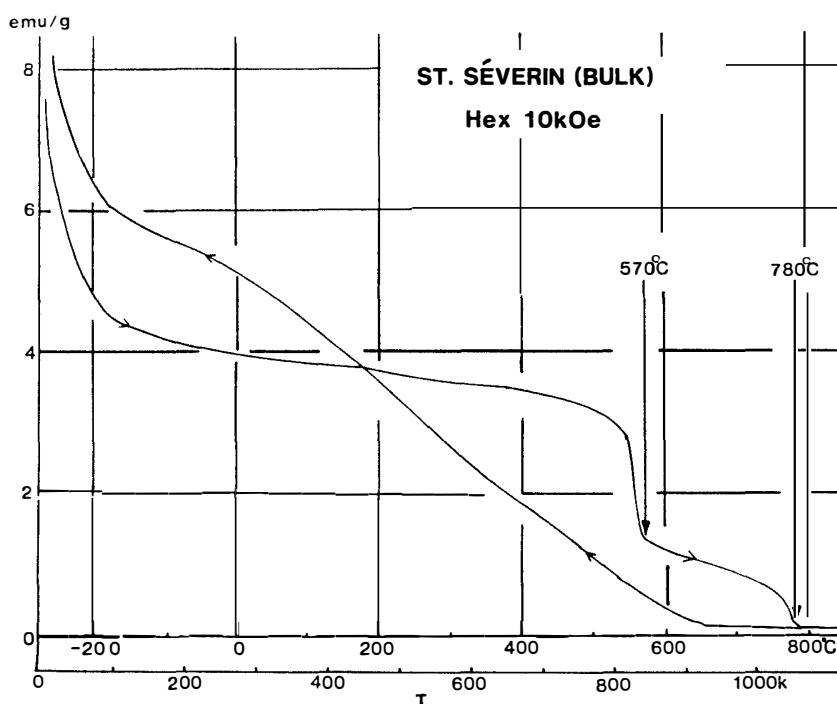


Fig. 1. First-run thermomagnetic curves of bulk specimen of St. Séverin LL6 chondrite.

Figures 2a–2d illustrate thermomagnetic curves of matrix, clast, whole metallic component and apparent tetrataenite phase extracted from metallic grains respectively, all in a magnetic field of 10 kOe for a temperature range between 20 and 820°C . In the first-run heating thermomagnetic curves of all bulk, matrix, clast, metal and extracted tetrataenite of St. Séverin, a sharp magnetic transition takes place at 560 – 565°C , which may correspond to Curie point (θ_c) of taenite of 56 wt% Ni. In the first-run heating thermomagnetic curves of bulk, matrix, clast and whole metal, a magnetic transition at 780 – 785°C corresponds to the $\alpha \rightarrow \gamma$ transition point ($\theta_{\alpha \rightarrow \gamma}^*$) of kamacite containing a small amount of cobalt. In the first-run cooling thermomagnetic curves of bulk, matrix and clast, the $\gamma \rightarrow \alpha$ transition point ($\theta_{\gamma \rightarrow \alpha}^*$) of kamacite phase is not clearly detected, but the $\theta_{\gamma \rightarrow \alpha}^*$ point of kamacite is clearly observable at 665°C in both first- and second-run cooling curves of whole metal and in the second-run cooling curve of clast. The $\gamma \rightarrow \alpha$ transition phenomenon in the first-run cooling curves of bulk, matrix and clast may be masked by magnetization of taenite phase

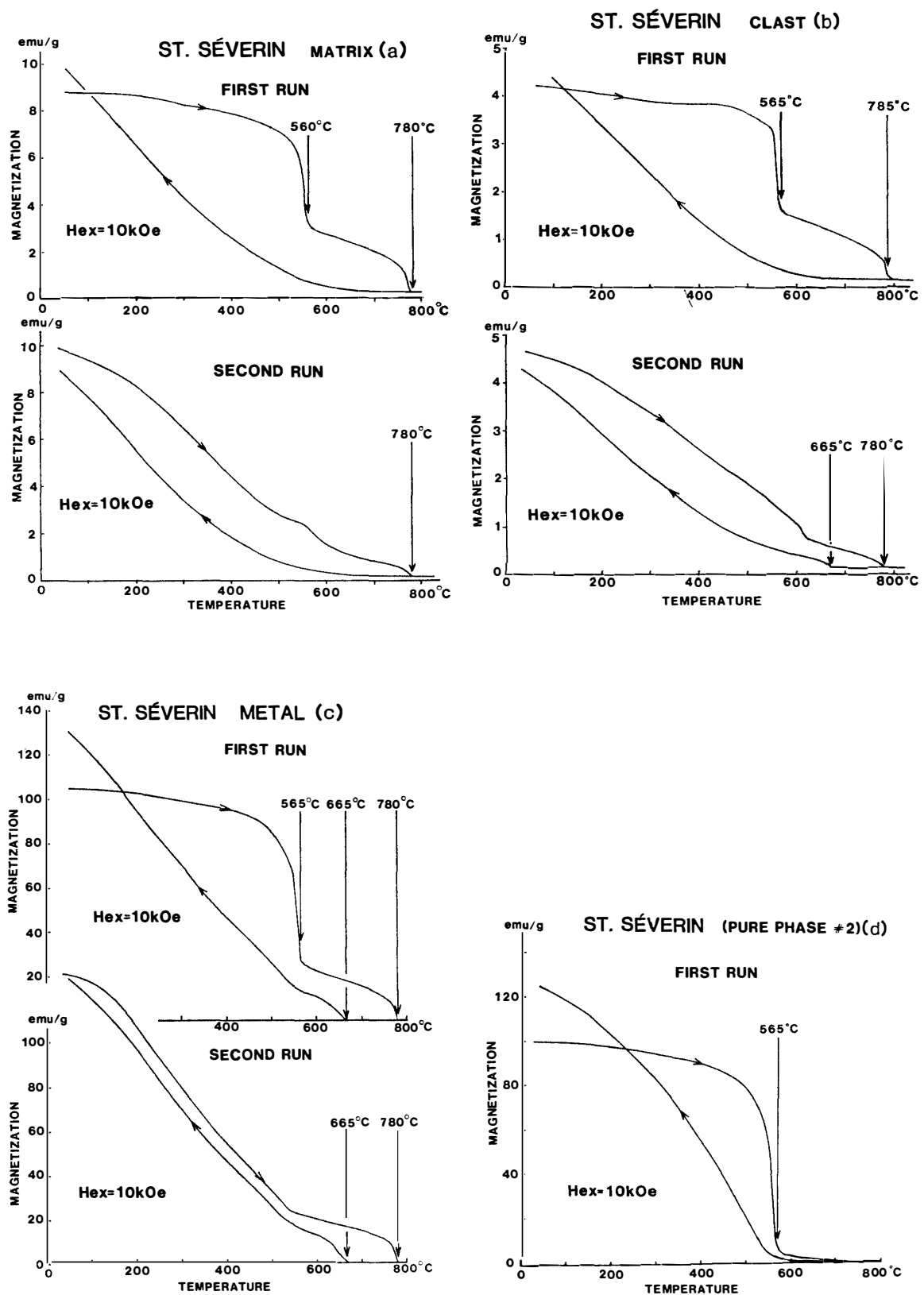


Fig. 2. First-run and second-run thermomagnetic curves of (a) matrix, (b) clast, (c) whole metal and (d) extracted apparent tetrataenite piece of St. Séverin.

which has a broad range of Curie points of a widely dispersed range in Ni content after the breakdown of tetrataenite phase at elevated temperatures. The kamacite phase of $\theta_{\alpha \rightarrow \gamma}^* = 780\text{--}785^\circ\text{C}$ and $\theta_{\gamma \rightarrow \alpha}^* = 665^\circ\text{C}$ corresponds to kamacite of 5.0 wt% Ni.

Figure 3 shows thermomagnetic curves of a man-made 50Fe50Ni alloy, which represents a disordered taenite of 50 wt% Ni, for reference. The thermomagnetic curves are approximately reversible with temperature and Curie point is 515°C . Comparing the thermomagnetic curves of the apparent tetrataenite phase (Fig. 2d) with Fig. 3, it will be noted (a) that the heating thermomagnetic curve of St. Séverin apparent tetrataenite phase decreases with temperature much less than that of 50Fe50Ni alloy in a temperature range lower than about 450°C , and (b) that the cooling thermomagnetic curve of St. Séverin apparent tetrataenite is a little less convex than that of 50Fe50Ni alloy, though the general features of the former look similar to those of the latter, and further (c) that the apparent Curie point of the cooling curve is lower than that of the heating curve for St. Séverin apparent tetrataenite phase.

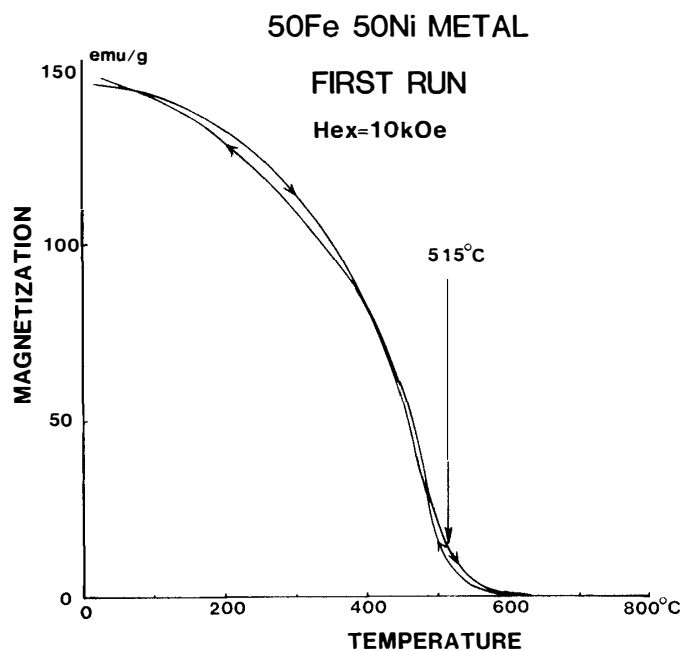


Fig. 3. Thermomagnetic curve of man-made 50Fe50Ni alloy.

The characteristic feature of thermomagnetic change of tetrataenite that the shape of its first heating thermomagnetic curve is essentially flat up to about 450°C and then drops abruptly to Curie point has already been discussed for Y-74160 LL7 chondrite (NAGATA and FUNAKI, 1982). As for St. Séverin LL6 chondrite, the first-run heating thermomagnetic curves of apparent tetrataenite component of matrix, clast and whole metal (*i.e.* magnetization of kamacite component being eliminated) are plotted together with the first-run heating thermomagnetic curve of the extracted apparent tetrataenite specimen itself in Fig. 4, where all thermomagnetic curves are normalized at 50°C . It may be concluded in Fig. 4 that the thermomagnetic characteristics of tetrataenite phase in matrix and clast of St. Séverin are substantially the same as those of extracted relatively large grains (11–53 mg in weight) of whole metal and apparent

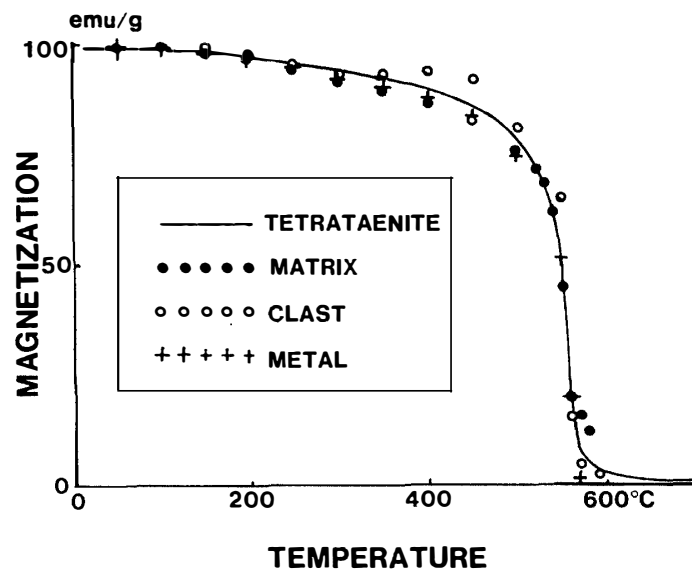


Fig. 4. Heating thermomagnetic curves of tetrataenite components in St. Séverin in matrix, clast and whole metal in comparison with apparent tetrataenite.

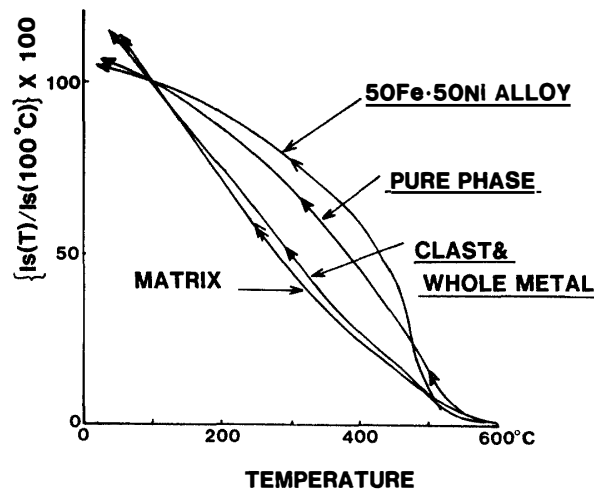


Fig. 5. Cooling thermomagnetic curves of tetrataenite components after breakdown of tetrataenite structure by heating in comparison with thermomagnetic curve of 50Fe50Ni alloy of disordered taenite structure.

tetrataenite metal in the first-run heating process.

On the other hand, general features of the first-run cooling thermomagnetic curves of taenite component (*i.e.* kamacite magnetization being eliminated) in matrix, clast and whole metal are considerably deviated from those of extracted apparent tetrataenite grain and 50Fe50Ni alloy, as shown in Fig. 5, where all thermomagnetic curves are normalized at 100°C. Namely, the cooling thermomagnetic curves of taenite components of matrix, clast and whole metal are concave, while those of extracted apparent tetrataenite and 50Fe50Ni alloy are convex. This result may suggest that the original ordered tetrataenite phase is not only broken down to transform to a disordered taenite, but tends to be homogenized with low-Ni taenite phases to form dis-

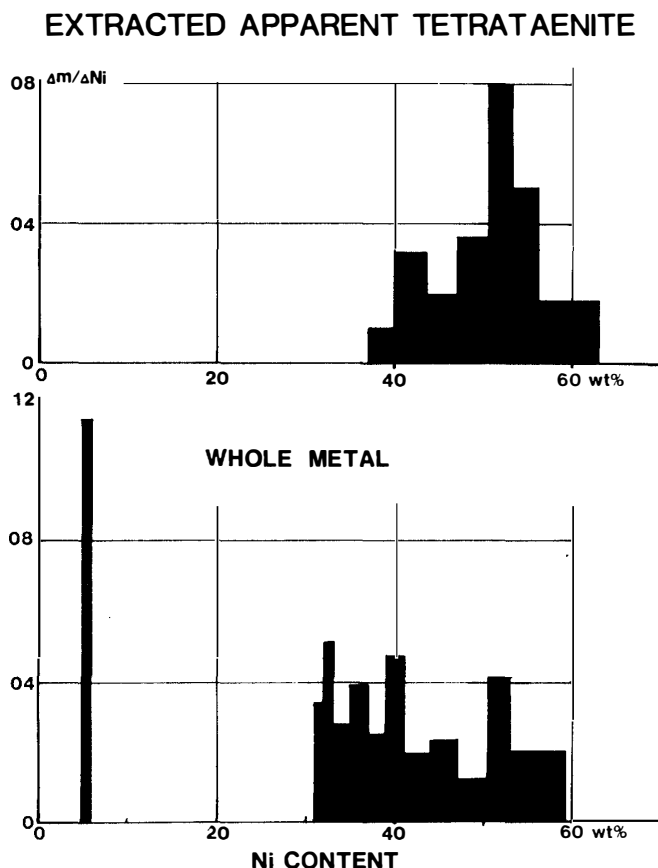


Fig. 6. Ni-content spectra for extracted tetrataenite piece (top) and whole metal piece (bottom) of St. Séverin.

ordered taenites of various Ni-contents of less than 50 wt%.

Figure 6 shows examples of Ni-content spectra of the extracted apparent tetrataenite sample (top) and the whole metal sample (bottom) after heating to 820°C, obtained by a thermomagnetic analysis of the first-run cooling thermomagnetic curve. Ni-content spectra of metallic component in matrix and clast of St. Séverin are approximately similar to the Ni-content spectrum of the whole metal. Namely, the Ni-content spectra of matrix, clast and whole metal of St. Séverin are spread over a wide range of Ni-content, extending to about 30 wt%Ni on the low-Ni side, in addition to a sharp spectral line of kamacite of 5 wt%Ni.

As already reported (DANON *et al.*, 1979a), the apparent tetrataenite phase in St. Séverin consists of genuine ordered tetrataenite crystallites (0.15–0.30 μm in diameter) and paramagnetic (less than 30 wt% in Ni-content) disordered taenite matrix according to results of Mössbauer spectral analysis. It seems most likely therefore that tetrataenite crystallites are homogenized with the surrounding low-Ni taenite to form the observed Ni-poorer disordered taenite phases of various Ni-content values at elevated temperatures. In the case of the extracted apparent tetrataenite sample, which is a relatively large grain (11 mg in weight) and whose surface is chemically cleaned up, almost no kamacite is detected by an Mössbauer spectral analysis so that the assumed homogenization process of Ni-content might be much less effective at

elevated temperatures.

The main interest in the present study is mostly concerned with a possible process of formation of tetrataenite phase from disordered taenites in meteorites in the extra-terrestrial space. It appears in the present study that a process of (tetrataenite)+(paramagnetic taenite) \rightarrow (disordered ferromagnetic taenite) is possible to take place at elevated temperatures. It seems possible than that a reverse process to form tetrataenite from disordered ferromagnetic taenites can take place on appropriate conditions. In the case of St. Séverin, however, the coexistence of kamacite phase also has to be taken into account.

2.4. Natural remanent magnetization

The natural remanent magnetization (NRM) of St. Séverin bulk sample is extremely stable against the AF-demagnetization, as shown by examples given in Figs. 7a and 7b, where the intensity and direction of residual NRM after AF-demagnetizing up to \tilde{H} (in Oe peak) are plotted. The residual NRM intensity after AF-demagnetizing up to 1800 Oe peak still keeps about 70% of the initial value, though some intensity fluctuations due to an inhomogeneity of NRM are involved in the AF-demagnetiza-

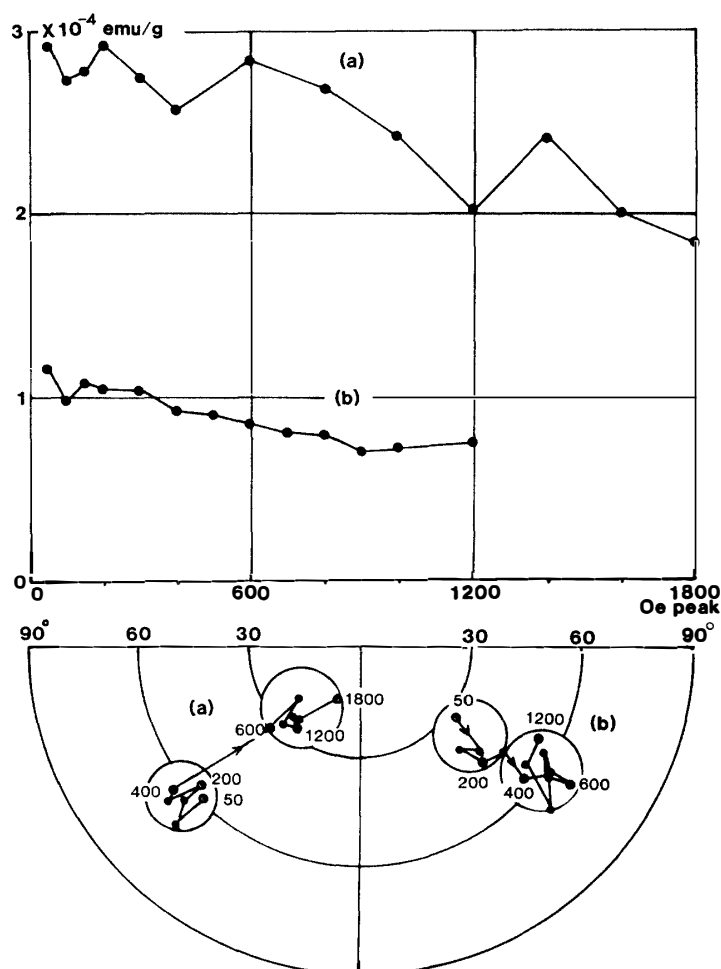


Fig. 7. AF-demagnetization curves of NRM of 2 bulk specimens of St. Séverin. (Top) Intensity change. (Bottom) Direction change.

tion curve. The residual NRM direction during the course of AF-demagnetization is one-directionally changed by about 40° in angle, but it is settled within 10° in angle around the mean direction for \tilde{H} values larger than 300–400 Oe peak, suggesting that St. Séverin meteorite possesses an extremely stable major component of NRM.

Figures 8 and 9, show thermal demagnetization behaviours in intensity and direc-

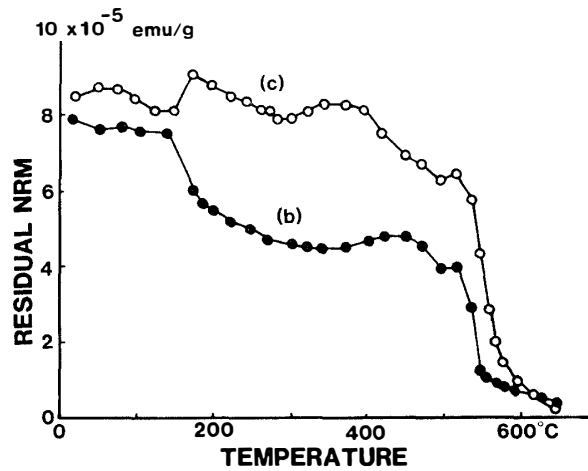


Fig. 8. Thermal demagnetization curves of NRM intensity of 2 bulk specimens of St. Séverin.

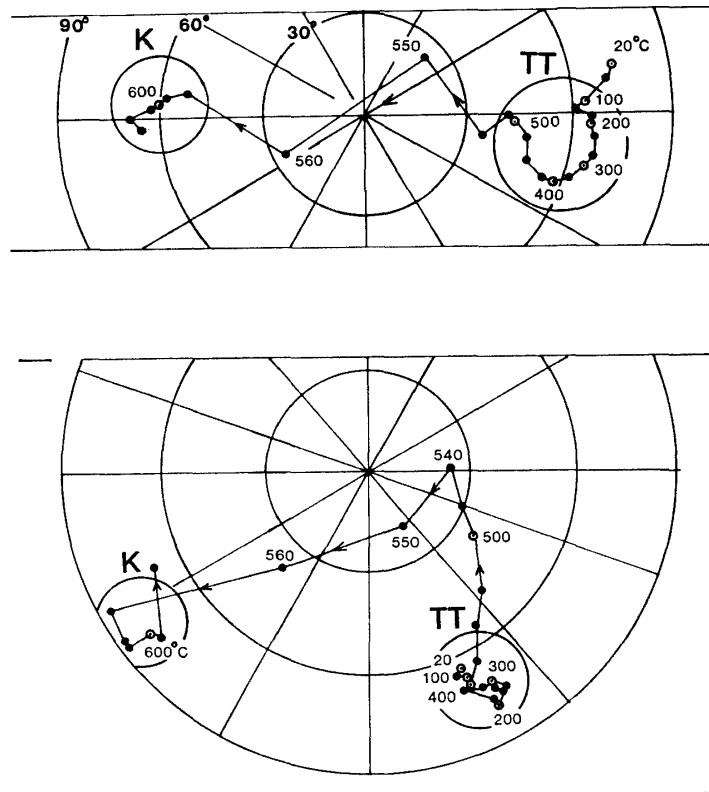


Fig. 9. Changes in direction during thermal demagnetization of NRM of 2 bulk specimens of St. Séverin. Specimen (b) (top) and specimen (c) (bottom).

tion respectively for the stable NRM component of St. Séverin bulk specimen after AF-demagnetizing up to 1200 Oe peak. In Fig. 8, the thermal demagnetization curves of NRM for two specimens, (b) and (c), are not simply monotonous but associated with some fluctuations, suggesting that NRM of St. Séverin meteorite consists of some heterogeneous components, probably caused by a variety of ordering degree of tetrataenite structure. Generally speaking, however, the extremely stable NRM component possessed by the tetrataenite component is thermally demagnetized almost completely between 500 and 560°C in temperature, a small portion of NRM still remaining at temperatures above 560°C. Figure 9 shows that the direction of NRM is kept approximately invariant for temperatures below 450–500°C during the course of thermal demagnetization, suggesting that the direction of NRM possessed by the tetrataenite component is reasonably stable within the temperature range. In the heating process above 500°C, the NRM direction changes largely up to almost 600°C, and then it is again stably settled at temperatures higher than 600°C, suggesting that a remaining NRM of a much smaller magnitude is possessed by the kamacite component at the elevated temperatures.

It may be generally concluded that NRM of St. Séverin meteorite is extremely stable as far as temperature is lower than about 400°C and the main parts of the stable NRM are possessed by the tetrataenite component. However, the acquisition process for the unusually stable NRM of tetrataenite phase has not yet been definitely identified.

3. Appley Bridge (LL6) and Tuxtuac (LL5) Chondrites

Both Appley Bridge and Tuxtuac are “falls” meteorites; Appley Bridge fell on England on October 13, 1914, and Tuxtuac fell on Mexico on October 16, 1975. WASILEWSKI (1982) has already experimentally demonstrated that Appley Bridge LL6 chondrite also shows the characteristic features of thermomagnetic curves of tetrataenite phase. In the present work, composition and structure of metallic components of Appley Bridge and Tuxtuac chondrites are first determined, and then magnetic properties of their bulk specimens are examined.

3.1. *Composition and structure of metal*

The average composition of Appley Bridge metal, determined with the aid of Mössbauer spectral analysis, is approximately given by

(Tetrataenite): (Disordered taenite): (Kamacite)=80: 20: (~0) (in wt%) .

The average composition of metals in Tuxtuac determined by the same Mössbauer spectral analysis is approximately given by

(Tetrataenite): (Disordered taenite): (Kamacite)=15: 85: (~0) (in wt%) .

In both chondrites, kamacite phase is hardly detectable in the Mössbauer spectral analysis, and the share ratios of ordered tetrataenite to disordered taenite in the two LL chondrites, are approximately reversed to each other.

3.2. *Basic magnetic properties*

Basic magnetic hysteresis curve parameters measured at room temperature and

Table 3. Basic magnetic properties of Appley Bridge (LL6) and Tuxtuac (LL5).

Sample	I_s (emu/g)		I_R (emu/g)		H_C (Oe)		H_{RC} (Oe)	
	Before Heating	After Heating	Before Heating	After Heating	Before Heating	After Heating	Before Heating	After Heating
Appley Bridge								
300 K	1.73	2.42	0.121	0.031	160	14.5	470	70
4.2 K	1.9	3.1	0.415	0.153	190	60	1210	3710
Tuxtuac								
300 K	2.93	3.1	0.055	0.015	40	11	255	175
4.2 K	3.6	4.45	0.176	0.128	90	60	3120	410

at liquid He temperature of bulk specimens of Appley Bridge and Tuxtuac LL chondrites before and after heating to 820°C are summarized in Table 3. Just as in the case of St. Séverin, H_C and I_R/I_s values of Appley Bridge are drastically reduced by the heat treatment, indicating a breakdown of the ordered tetrataenite structure to the disordered taenite by the heat treatment. The decreasing changes of H_C and I_R/I_s values of Tuxtuac caused by the heat treatment are considerably smaller than those of Appley Bridge, indicating that the share ratio of tetrataenite phase in Tuxtuac metal is much smaller than that in Appley Bridge metal.

3.3. Thermomagnetic curve characteristics

Figures 10 and 11 show the first-run and second-run thermomagnetic curves of

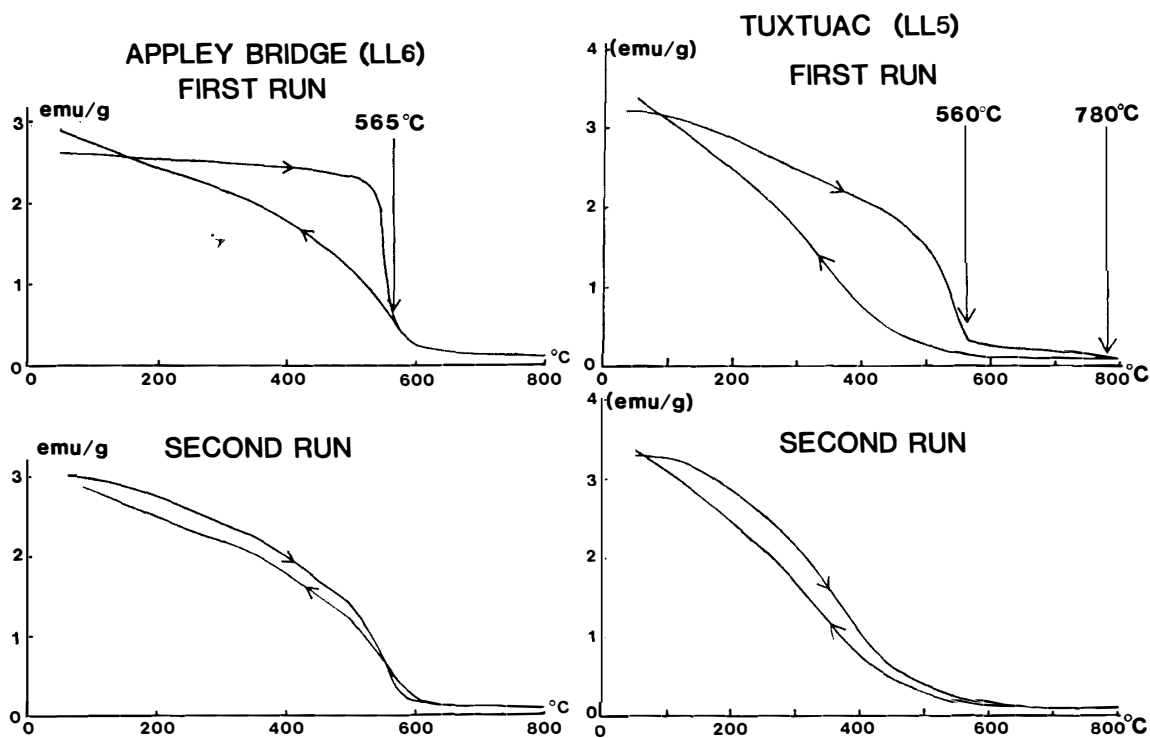


Fig. 10. First-run and second-run thermomagnetic curves of Appley Bridge LL6 chondrite.

Fig. 11. First-run and second-run thermomagnetic curves of Tuxtuac LL5 chondrite.

Appley Bridge and Tuxtuac respectively. As Curie point is 565°C and no kamacite phase is detected in Appley Bridge thermomagnetic curves, its ferromagnetic metallic component may consist of tetrataenite and disordered taenite of about 50%Ni in the original state. Actually, the first-run heating thermomagnetic curve has the characteristic flatness of tetrataenite for a temperature range up to 450°C , and the cooling thermomagnetic curves after the breakdown of tetrataenite phase by the heat treatment are approximately identical, in shape, to the cooling thermomagnetic curve of 50Fe50Ni alloy shown in Fig. 3.

In the Tuxtuac thermomagnetic curves, the characteristic flatness of the first-run heating thermomagnetic curve for tetrataenite phase appears to be superimposed on an ordinary thermomagnetic curve of disordered taenite phase, because relative content of tetrataenite phase is only 3/17 of disordered taenite phase.

Although a small amount of kamacite phase of $\theta_{\alpha \rightarrow \gamma}^* = 780^{\circ}\text{C}$ can be detected in the first-run heating thermomagnetic curve of Tuxtuac, it seems very likely that the kamacite phase is much reduced to a too small amount to be magnetically detected in the second-run heating curve. The first-run cooling thermomagnetic curve and the second-run curves of Tuxtuac are concave in a temperature range from 600 to 300°C and are essentially different from the convex cooling thermomagnetic curve of Appley Bridge as well as 50Fe50Ni alloy. With the aid of the same analysis method for St. Séverin, the Ni-content spectra of metallic component after the heat treatment for

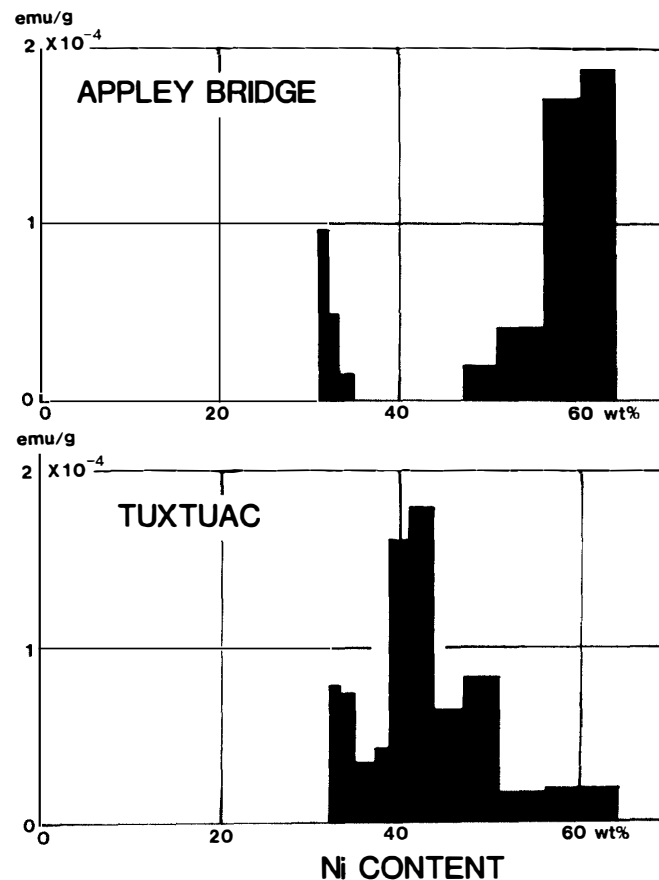


Fig. 12. Ni-content spectra in metallic component for Appley Bridge (top) and Tuxtuac (bottom).

Appley Bridge and Tuxtuac are obtained as shown in Fig. 12. As indicated in Fig. 10, the main components of metals after the heat treatment in Appley Bridge are disordered taenites of 50–60 wt%Ni, though a small amount of Ni-poor taenite of about 30 wt%Ni still coexists. As indicated in Fig. 11, on the other hand, the metallic components in Tuxtuac after the heat treatment are represented by a broad continuous band of Ni-content in taenite extending from 30 wt%Ni to 60 wt%Ni and having a maximum peak around 40 wt%Ni. No spectral line of kamacite phase is detectable in the Ni-content spectrum of Tuxtuac after the heat treatment, though about 7 wt% of kamacite phase of 5 wt%Ni in metal is magnetically detected in the first-run heating thermomagnetic curve. Only a possible interpretation of the observed disappearance of kamacite phase by heating will be a homogenization of kamacite phase with neighbouring taenite phases.

4. Thermomagnetic Characteristics of Tetrataenite-rich Meteorites

Thermomagnetic behaviours of tetrataenite phase can be characterized by several measurable factors. One of the characteristic factors will be a sharp irreversible magnetic transition in the first-run heating thermomagnetic curve, examples of which are typically represented by Fig. 2d for an extracted piece of apparent tetrataenite of St. Séverin and by Fig. 10 for Appley Bridge chondrite. The magnetic transition temperature, which is considered as Curie point of tetrataenite phase, is ranged between 560 and 570°C as summarized in Table 4, where observed values of $\theta_{\alpha \rightarrow \gamma}^*$ and

Table 4. Magnetic transition temperatures of tetrataenite-rich meteorites.

Meteorite	Tetrataenite θ_c (°C)	Kamacite	
		$\theta_{\alpha \rightarrow \gamma}^*$ (°C)	$\theta_{\gamma \rightarrow \alpha}^*$
Y-74160 (LL7)	560	None	None
Appley Bridge (LL6)	565	None	None
St. Séverin (LL6) (Bulk)	565	780	665
ALH-77260 (L3)	570	780	600–700
Tuxtuac (LL5)	560	780	—
ALH-77219 (mesosiderite)	560	760	630

$\theta_{\gamma \rightarrow \alpha}^*$ of coexisting kamacite phase, if any, also are given. In the cases of Y-74160 and Appley Bridge, their second-run thermomagnetic curves clearly show a single Curie point of the transformed disordered taenite of about 55 wt%Ni around 550°C, so that it will be highly plausible that their sharp magnetic transition point at 560–565°C in the first-run heating thermomagnetic curve corresponds to Curie point (H_c) of tetrataenite phase. Since, however, Curie points of Fe-Ni alloy at 560° and 570° correspond respectively to 57 and 59 wt% in Ni-content on the basis of a Curie point vs. Ni-content relationship for taenite given by CRANGLE and HALLAM (1963) and an expected Curie point of a stoichiometric tetrataenite composition (*i.e.* 48.8 wt%Fe, 51.2 wt%Ni) will be about 505°C, there will be a possibility that Curie point of the ordered tetrataenite is a little higher than Curie point of disordered taenite of the same chemical composition.

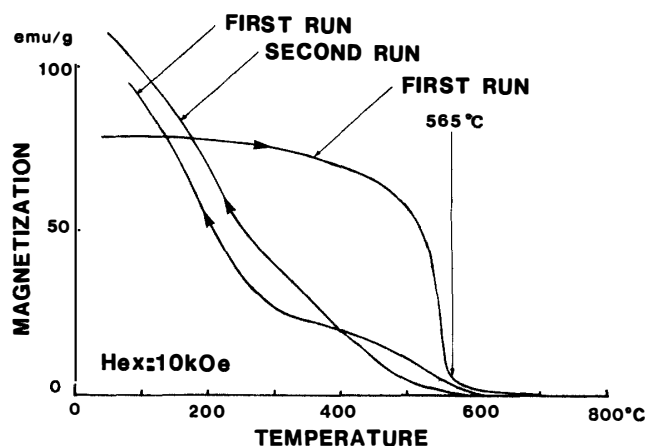


Fig. 13. First-run thermomagnetic curves and second-run cooling thermomagnetic curve of Santa Catharina Ni-rich ataxite.

In order to ascertain the sharp irreversible magnetic transition phenomenon of tetrataenite-rich metal, magnetic characteristics of Santa Catharina Ni-rich ataxite, which has been crystallographically studied in detail to prove that the ordered tetrataenite is its main constituent (DANON *et al.*, 1979b), are examined as much precisely as possible. The chemical composition of tetrataenite phase in Santa Catharina is (48–49) wt%Fe (50–51) wt%Ni, while the characteristic sharp magnetic transition point temperature (θ_c) is 565°C, as shown in Fig. 13. Taking into consideration thermomagnetic characteristics of an extracted piece of apparent tetrataenite phase of St. Séverin shown in Fig. 2d also, it may be concluded that the ordered tetrataenite has a sharp magnetic transition point at 560–565°C, and it may be Curie point of the ordered tetrataenite structure.

A similar sharp magnetic transition at 560–570°C observed in the other meteorites, in which kamacite phase coexists, can be interpreted in the same way.

The second point may be concerned with characteristics of the cooling thermomagnetic curve after the breakdown of the ordered tetrataenite structure by heating. In the cases of Y-74160 and Appley Bridge, their cooling thermomagnetic curves may represent mostly the reversible thermomagnetic curve of disordered taenite of 50–55 wt% in Ni-content. However, the original values of magnetic coercive force (H_c) at room temperature of Y-74160 and Appley Bridge are only 255 Oe (NAGATA and FUNAKI, 1982) and 160 Oe (Table 3) respectively, which are much smaller than the expected H_c values of tetrataenite itself. In fact, the observed H_c value of Santa Catharina Ni-rich ataxite at room temperature amounts to 2800 Oe. Since the magnetic coercive force (H_c) of a tetrataenite crystal along its uniaxial easy axis is $H_c=4900$ Oe, the observed H_c value of Santa Catharina is reasonable for an assemblage of almost randomly oriented tetrataenite microcrystallites.

As already discussed (NAGATA and FUNAKI, 1982), the magnetic coercive force of a mixture of a magnetically hard component having a large coercive force and a soft component having a small coercive force is affected more largely by the soft component than by the hard component, thus resulting in a much smaller resultant coercive

force than the coercive force of the hard component. Assuming then that the magnetic coercive force of an assemblage of randomly oriented tetrataenite microcrystallites is 2800 Oe, weight content (m) of tetrataenite phase in metallic components of Y-74160 and Appley Bridge are estimated as $m=43$ wt% for Y-74160 and $m=67$ wt% for Appley Bridge where the soft components are disordered taenite of $H_c=8$ Oe for Y-74160 and disordered taenite of $H_c=4.5$ Oe for Appley Bridge. It is observed by an electron microprobe analysis that the chemical composition of disordered taenite phase is about 50 wt%Ni in Y-74160. Therefore, the first-run cooling curve after the breakdown of tetrataenite structure shows a typical convex curve which is very similar to the thermomagnetic curves of 50Fe50Ni alloy (Fig. 3), and the Ni-content spectra corresponding to these thermomagnetic curves consist of spectral lines concentrated around 50%Ni. Thermomagnetic curve characteristics and the corresponding Ni-content spectra for Appley Bridge are very similar to those of Y-74160, so that the disordered taenite phase of Appley Bridge also will be around 50%Ni in chemical composition. Thermomagnetic characteristics of the apparent tetrataenite piece extracted from St. Séverin metal also show similar characteristics, probably because the chemical composition of remaining disordered taenite is near 50%Ni in composition, as suggested by Fig. 6.

On the other hand, Ni-content spectra of taenite component in the whole metal of St. Séverin (Fig. 6, bottom) and in Tuxtuac (Fig. 12, bottom) are spread over a fairly wide range of Ni-content (30–60 wt%) after the breakdown of tetrataenite phase by heating. As already discussed individually for the cases of St. Séverin and Tuxtuac, it seems very likely that fine ordered tetrataenite specks of 50Fe50Ni in atomic composition are not simply broken down to the disordered taenite structure but also tend to be homogenized with surrounding low-Ni taenites, and probably even with kamacite, at elevated temperatures. This hypothesis of homogenization of Ni-content in the disordered taenite (namely, f.c.c. structure of Fe-Ni alloy) after the breakdown of tetrataenite structure at elevated temperatures may need further experimental certifications with respect to microstructure of the metallic grains concerned before and after the heat treatment. It seems, however, that an example of thermomagnetic characteristics of Santa Catharina Ni-rich ataxite, shown in Fig. 13, may support the hypothesis of homogenization of Ni-content. In Fig. 13, the first-run heating and cooling thermomagnetic curves and the second-run cooling thermomagnetic curve of Santa Catharina are shown. The second-run heating thermomagnetic curve is almost identical to the first-run cooling curve, and the third-run heating and cooling thermomagnetic curves are almost same as the second-run cooling curve. Namely, the first-run cooling and second-run heating thermomagnetic curves after the first heat treatment show that the ferromagnetic metal consists of two main components, a component having Curie point at 500–550°C (corresponding to 50–55 wt%Ni) and the other component having Curie point at 250–300°C (corresponding to 35–40 wt%Ni), while the second-run cooling and the third-run heating and cooling thermomagnetic curves after the second and third heat treatments show that the ferromagnetic metal consists of a continuous spectra of Ni content in a range of 30–55 wt%Ni, as in the cases of St. Séverin whole metal and Tuxtuac.

5. Concluding Remarks

In the present study, magnetic properties of Appley Bridge, Tuxtuac and St. Séverin tetrataenite-rich chondrites are examined in comparison with Y-74160 tetrataenite-rich chondrite, Santa Catharina Ni-rich ataxite and an artificially made 50Fe50Ni alloy. In particular, bulk specimens, matrix, clast, metallic grains and an extracted piece of apparent tetrataenite phase of St. Séverin are separately examined in some detail.

Ferromagnetic features of tetrataenite component in meteorites are characterized by (a) a high magnetic coercivity represented by unusually large values of H_C and H_{RC} , (b) an essential flatness of thermomagnetic curve in a temperature range below about 450°C, as shown in Fig. 4, and (c) a breakdown of these typical magnetic characters of tetrataenite structure by heating up to temperature above Curie point during a relatively short period, *i.e.*, an hour in order of magnitude. Because of the high magnetic coercivity of tetrataenite component, NRM of tetrataenite-rich meteorite contains an extremely stable component which is hardly AF-demagnetized even by 2000 Oe in alternating magnetic field intensity, as demonstrated for Y-74160, ALH-77260 and St. Séverin chondrites.

However, the formation process for the tetrataenite phase in meteorites and the acquisition mechanism for the stable NRM of tetrataenite-rich meteorites have not yet been definitely clarified. As each ordered tetrataenite phase in meteoritic metals occupies only a very fine volume of several μm or less in mean diameter in matrix of disordered taenite, microscopic examinations, both crystallographically and magnetically, of tetrataenite phases in meteoritic metals in much more detail will be required in further studies.

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