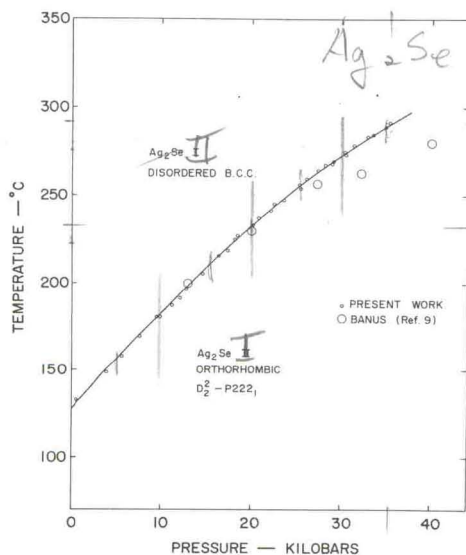
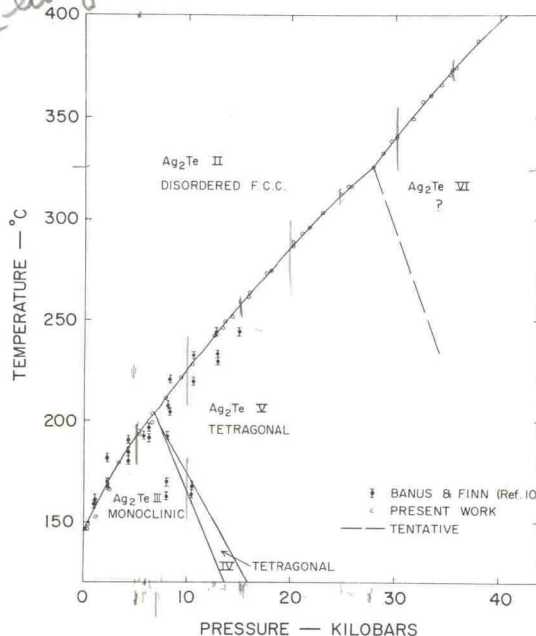
Fig. 2. Phase diagram of Ag_2S .Fig. 3. Phase diagram of Ag_2Se to 300°C and 40 kbars (including the work of Banus).Fig. 4. Phase diagram to 400°C and 40 kbars of Ag_2Te (including the work by Banus and Finn).

of the II/III/IV triple point is the one at 12.3 kbar, 195°C . However, its location is not on the extrapolation of Bridgman's III/IV boundary. Unfortunately we did not have a sufficient amount of sample of high purity to repeat Bridgman's volumetric work.

Phase relations in the system silver-sulfur

and the transitions in silver sulfide were investigated by Kracek[6] and reviewed by Frueh[21]. The temperature-composition phase diagram reveals that the III/II transi-

*Enlarge

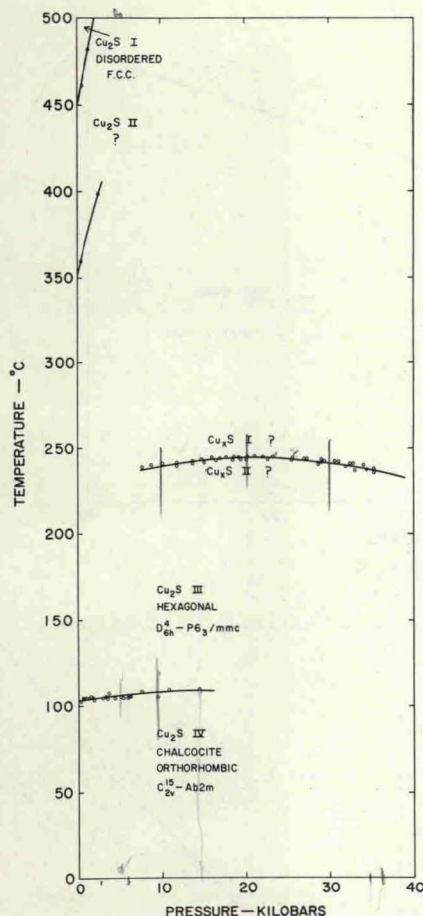


Fig. 5. Phase diagram of Cu_2S with the phase boundary Cu_xS included.

tion temperature on the silver-rich side is slightly lower than that on the sulfur-rich side ($176.3^\circ \pm 0.5^\circ\text{C}$ as compared with $177.8^\circ \pm 0.7^\circ\text{C}$). Wagner [22], from EMF measurements on galvanic cells containing solid Ag_2S , concluded that Ag_2S in equilibrium with metallic silver at 200°C contains 2×10^{-3} g atom Ag/mole Ag_2S excess silver whereas Ag_2S in equilibrium with sulfur has nearly the ideal stoichiometric composition. At 160°C the variability in stoichiometry is 3×10^{-5} g atom Ag/mole Ag_2S . He made a thermodynamic calculation to account for the 1.7°C difference between transition temperatures on the Ag-rich and S-rich side. Aside from this small

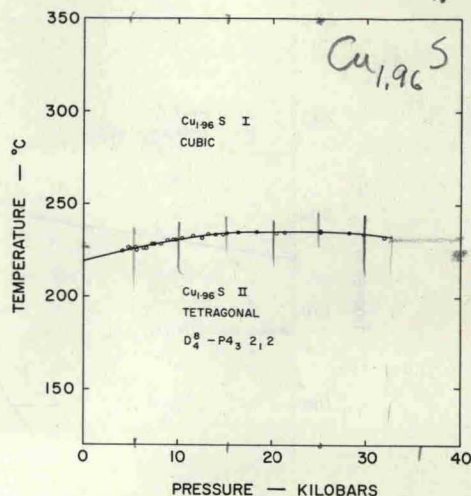


Fig. 6. The I/II phase boundary in $\text{Cu}_{1.96}\text{S}$.

difference the transition temperature does not vary further with excess silver or sulfur, and we consider it safe to have carried out the present experiments on Ag_2S enclosed in silver capsules. The II/I transition is much more sensitive to excess sulfur or silver [6]. We made several attempts to investigate the course of the II/I transition with pressure but encountered a vigorous endothermic sample-capsule reaction above 500°C . This was the only case that such a reaction was encountered, possibly a dissolution of silver in the Ag_2S sample. No such reactions were encountered in the other silver and cuprous chalcogenides at lower temperatures.

Silver selenide

The α/β transformation [8] in Ag_2Se had been studied previously by Roy *et al.* [7] and by Banus [9]. In our experiments we first searched for DTA signals around 130°C , corresponding to the α/β transition. At first no such signals could be obtained, but we found that if the sample was heated in situ to $\sim 350^\circ\text{C}$ for a period of several minutes, sharp DTA signals were obtained after the sample had been cooled and another search was made around 130°C . Such signals were then obtained repeatedly four to five times,