Garnet-Pyroxene Alteration Mapping in the Ludwig Skarn (Yerington, Nevada) with Geoscan Airborne Multispectral Data*

Abstract
Geoscan airborne multispectral image data of the skarn and calc-silicate-metamorphosed rocks near Ludwig (Yerington district, Nevada) have been analyzed and compared with published and unpublished maps. This study examines the spectral differentiation between two stages of garnet-pyroxene alteration, early metamorphic skarn and late metasomatic skarn. Both types of calc-silicate rock were successfully delineated using thermal infrared band difference images. Differentiation between the two alteration styles was achieved using a (0.717-μm - 0.873-μm) difference image, based on the deeper 0.87-μm Fe³⁺ garnet absorption in skarn.

Compositional variations, grain size, and weathering styles were examined as possible sources of this spectral variation. While all three contribute, weathering characteristics have the greatest effect. Skarnoid contains finer-grained and greater proportions of pyroxenes than skarn, which readily weather to a limonitic coat that mutes the 0.87-μm absorption.

Introduction
In their review paper on skarn deposits, Einaudi et al. (1981) describe skarn as "coarse-grained Ca-Fe-Mg-Mn silicates formed by replacement of carbonate-bearing rocks accompanying regional or contact metamorphism and metasomatism." The majority of the world's tungsten is produced from skarn deposits and this deposit type is an important source of copper, iron, molybdenum, and zinc.

Despite their economic value, there is a distinct paucity of studies applying remote sensing to skarn deposits. The vast majority of remote sensing efforts in economic geology have been directed toward the identification of iron-staining or hydroxyl-bearing alteration products in porphyry copper or epithermal districts, or in the determination of regional structural trends. Skarns are a common feature in carbonate-bearing strata intruded by porphyry copper plutons (Einaudi, 1982) but, as Abrams and Brown (1985) comment in a study of the Silver Bell district, Arizona, "detection of most tactite (skarn) is unlikely because of the small size of outcrops and the lack of spectrally diagnostic minerals."

The increased spatial and spectral resolution of modern imaging spectrometers, however, suggests that such assertions be reevaluated. The Ludwig skarn in the Yerington district, Nevada, was selected for such a study. The geology of the area is well-characterized (Harris and Einaudi, 1982; Proffett and Dilles, 1984; Dilles and Einaudi, 1992), while extensive exposures and sparse vegetation cover provide ideal conditions for remote sensing. This study employs Geoscan AMSS Mk II image data and follows an approach similar to that of an exploration program:

- description of the setting and the target alteration types,
- selection of empirical image treatments based on spectral characteristics of the dominant minerals (acquired from both lab measurements of reconnaissance field samples and from the literature),
- comparison of the image processing results with geologic maps, and
- identification of the sources of spectral differentiation to determine whether the treatments can be exported to similar targets or are limited to the conditions present in the study area.

Instrumentation and Methods
The Geoscan AMSS Mk II scanner was first described by Lyon and Honey (1990). The AMSS Mk II was developed as a tool for mineral exploration by Geoscan Pty. Ltd., a division of Carr-Boyd Minerals based in Perth, West Australia. It is an imaging spectrometer using grating-dispersive optics with three sets of linear array detectors, one each in the visible/near infrared (VNIR), short-wave infrared (SWIR), and thermal infrared (TIR) (Table 1). The data set used in this study was flown 21 July 1990 (1250 PDT) at an elevation of 4700 ft above ground level (AGL), with a nominal pixel resolution of 3m.

Data acquisition by the Geoscan system differs from most other scanning systems in that the instrument gains and offsets are not fixed, but are adjusted for the conditions present along the flight line. The system is flown twice over the target area. During the first flight, an operator adjusts the gains and offsets in each channel to maximize the surface contrast into an 8-bit dynamic range (0 to 255). The same area is then flown a second time with the channels held constant at the new gains and offsets. By adjusting the offset values such that all channel means have a digital number (DN) of 127, the Geoscan data are effectively pre-corrected for the solar radiation curve (Rubin, 1991; 1993). This lessens the necessity for calibrations such as the "flat field" and log-residual (Green and Craig, 1985; Roberts et al., 1986) in order to compare image data with reflectance. This method does require corrections for variation in the offset and gain settings (Windeler, 1992); raw Geoscan data were transformed to apparent reflectance using the "two-point correction" method.


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method of Lyon et al. (1975). Examples of transformed visible/near-infrared data are provided in the section on Image Data and Treatments. No thermal infrared spectrometer was available for this study, however, and thus uncalibrated thermal image data were used.

The data set exhibited considerable distortion perpendicular to the east-west flight direction, even after applying a panoramic correction. This distortion resulted from the combination of a relatively low flight level with considerable topographic relief (400-m increase from west to east on the image). The image data were registered to a 1:4800-scale topographic map using drainage junctions or other distinct geographic features as control points. The average RMS error between the registered image and the map is 7 to 8 pixels.

Color composite images in this study are described with a shorthand notation. The three single bands or band differences in an image are each given a subscript denoting their screen display color. For example, a “true color” treatment would be described as (0.522 µm)g(0.583 µm)b(0.645 µm)r or, in Geoscan channels, 1g 2b 3r.

Over 350 reflectance spectra were acquired from 202 samples collected by the author and 37 samples provided by M.T. Einaudi. Spectra were recorded from 0.4 to 2.5 µm, using an IRIS Mark IV spectroradiometer manufactured by Geophysical Environmental Research, Inc. Halon was used as a reflectance standard.

Garnet and/or pyroxene compositions were analyzed by electron microprobe in 23 thin sections. These analyses were performed on a JEOL Superprobe 733 and calculated using ZAF corrections on a Kevex Sesame-Delta system. Typical operating conditions were 15-keV accelerating voltage, 20-nA beam current, 10-µm spot size, and 30-second count time. Albite was used as an Al and Si standard. Wollastonite and spessartine were standards for Ca and Mn, respectively, while oxides were used for Fe, Mg, and Ti.

**Geology**

Numerous workers have contributed to the understanding of the Yerington district, extending back to Knopf (1918). Regional studies have been conducted by Proffett (1977) and Proffett and Dilles (1984; in press). Information regarding the petrology and alteration of the Yerington batholith are provided by Dilles (1987) and Dilles and Einaudi (1992). Descriptions of skarn mineralogy and genesis in the district are summarized from Einaudi (1977), Harris (1979), and Harris and Einaudi (1982).

Triassic and Jurassic sedimentary and volcanioclastic rocks are exposed in the Singate Range near Yerington, Nevada (Figure 1). These Mesozoic rocks were intruded and altered by a Jurassic quartz monzodioritic to granodioritic batholith. Early stage alteration of the Mesozoic rocks was largely metamorphic, resulting in the formation of fine-grained, garnet-pyroxene hornfels (skarnoid) in argillite, volcanioclastic, and silty limestone units. Skarnoid is iron-poor, with intermediate grossular-andradite garnets and diopsidic pyroxenes. Early stage alteration also formed garnet-pyroxene and plagioclase-Idocrase-clinozoisite endoskarn in apophyses of quartz monzodiorite. Intrusion of granite porphyry dikes was accompanied by porphyry copper mineralization in the batholith and skarn formation in a massive limestone unit. Late stage skarn is metasomatic and iron-rich; garnets commonly approach pure andradite, and pyroxenes are salitic. Tertiary Basin and Range faulting has rotated the district ~90° westward, resulting in the horizontal exposure of a cross-section through the district as it was in Jurassic time (Proffett, 1977).

The textural, compositional, and genetic features of hornfels and skarn in the study area are summarized and compared in Table 2. From an exploration standpoint, the distinction between these two mineralogically similar alteration types can be critical; at Yerington, early hornfels altera-
Hornfels
- early, mostly metamorphic
- extensively replaced silty limestone, calcareous argillite units; also replaced massive limestone and volcanoclastic units close to the igneous contact
- fine-grained (variable, most <0.5 mm)
- iron poor garnets 40-60 mol % andradite pyroxenes < 15 mol % hedenbergite
- barren, no associated mineralization

Skarn
- late, metasomatic
- replaced massive limestone unit
- coarse-grained (0.1 to > 10 mm)
- iron-rich garnets 65-100 mol % andradite pyroxenes 15-45 mol % hedenbergite
- mineralized, several small chalcopyrite orebodies

**Image Data and Treatments**

**Calc-Silicate Discrimination**

The region of the electromagnetic spectrum between 8 and 13 \(\mu\)m is of particular interest to geologists because of the fundamental vibration of the Si-O tetrahedral bond. This vibrational absorption is a key feature in the spectra of the silicate rocks that make up much of the Earth's crust. The location of this absorption has been shown to shift to progressively longer wavelengths with depolymerization of Si-O tetrahedra and substitution of atomically heavier cations for Si (Lyon, 1962; 1965). This absorption shifts from 9 \(\mu\)m in felsic rocks (quartz, K-and Na-feldspar-rich) to nearly 72 \(\mu\)m in the most mafic rocks (Ca, Fe, Mg-rich) and thus can be an important tool for rock type identification.

Emittance spectra for quartz, feldspar, epidote, and grossular garnet are presented in Figure 2. The emittance minima for quartz and garnet are separated by more than 2 \(\mu\)m, a fortuitous occurrence that aids in discrimination between the two. In terms of Geoscan thermal bands, quartz and quartz-feldspathic rocks will be relatively dark in the 8.64-\(\mu\)m and 9.17-\(\mu\)m channels (19 and 20), whereas very mafic rocks will appear relatively dark in the 10.22-\(\mu\)m and 10.75-\(\mu\)m channels (22 and 23). In the case of the skarn body, a relative gradient between these channels (determined by either difference or ratio) should thus indicate quartz-rich (positive), calc-silicate (negative), or intermediate (flat) lithologies.

In order to discriminate between garnet-pyroxene-altered rocks and other lithologies, a thermal band difference treatment has been applied to the Ludwig 3-m data set in Plate 1a. This image displays (10.75-\(\mu\)m - 9.70-\(\mu\)m)\(_a\), (10.75-\(\mu\)m - 9.17-\(\mu\)m)\(_c\), (10.75-\(\mu\)m - 8.64-\(\mu\)m)\(_r\), and has been smoothed using a 3 by 3 low pass filter to eliminate excess noise. In this orientation, north is to the right and the top of the image corresponds to exposures closer to the paleosurface. Garnet-pyroxene rock appears dark, while quartz and feldspar-rich material is bright. This treatment also identifies the drainages which carry alluvial skarn and hornfels. Hereafter, this difference treatment will be referred to by its Geoscan channels: (23-21)\(_a\), (23-20)\(_c\), (23-19)\(_r\).

A distribution map of garnet-pyroxene rock is presented in Plate 1b. Comparison between the (23-21)\(_a\), (23-20)\(_c\), (23-19)\(_r\) image and the geologic map shows an extraordinary correlation. It is apparent that the overall distribution and morphology of garnet-pyroxene skarn and hornfels are clearly defined by the thermal differences treatment. Several common points of reference are indicated by numbers 1 to 5 on Plates 1a and 1b; other features of note or mismatch are indicated on the image with smaller numbers. All annotated features are briefly described in Table 3.

Epidote also appears dark on this treatment because of its long-wavelength absorptions between 10.5 and 11 \(\mu\)m (Salisbury et al., 1991). This is most apparent near the southeastern edge of the image (area 6), where several dark anomalies not identified on the garnet-pyroxene map result from exposures of andesite altered to massive epidote.

Bright responses on the (23-21)\(_a\), (23-20)\(_c\), (23-19)\(_r\) image correspond to quartz- and feldspar-rich lithologies. The lone bright yellow spot on the northwest edge is a quartzite (area 7), while the north-northeast-trending white area immediately west of the main exposures of garnet-pyroxene rock results from a felsite unit (area 8). The bright area on the eastern edge is a tightly folded rhyolite flow (area 9).

**Figure 2.** Emittance spectra of quartz, feldspar, epidote, and grossular garnet. The quartz, feldspar, and epidote and grossular garnet spectra are direct emittance measurements from Lyon and Green (1975). The epidote spectrum is calculated from a reflectance measurement from Salisbury et al. (1991; sample Epidote.1, fracture surface) using Kirchoff's Law (\(E = 1 - R\)).
Gypsum dumps in the northwest (area 10) appear red to orange on the (23-21)R(23-20)R(23-19)R treatment. This was not initially expected but follows from the thermal spectrum of gypsum. The sulfate ion has intense absorptions at 8.3 and 8.7 μm which are narrow relative to the Si-O absorption in quartz (Lyon, 1984). This absorption would appear in the 8.64-μm and (to a lesser degree) 9.17-μm channels (19 and 20), and is thus brightest on the (23-19) difference (red on this treatment).

**Skarn/Hornfels Discrimination**

**Visable/Near-Infrared Spectral Features of Garnet-Pyroxene Skarn and Hornfels**

Thermal difference images were effective at defining areas of calc-silicate alteration in the study area, but could not differentiate between skarn and hornfels. The VNIR laboratory spectra of skarn and hornfels samples from Yerington were compared in order to identify features that could be used for discrimination. Garnet and pyroxene comprise the dominant mineralogy of these alteration types and generally determine the spectral features. Garnets in the study area fall within the grossular (Ca3Al2Si3O12) - andradite (Ca3Fe2Si2O12) series (Harris, 1979). With these compositions, the spectra of Yerington garnets are dominated by effects of Fe3+ in the octahedral site. The reflectance spectra of andradite-grossular garnets show two major Fe features: a strong drop in reflectance toward the UV due to charge transfer and a crystal field reflectance minimum at 0.87 μm (Hunt et al., 1973).

The visible and near-infrared spectral features of pyroxenes have been studied in much greater detail than those of the garnets, due to their probable importance in the remote sensing of planetary surfaces. A paper by Straub et al. (1991) contains a concise discussion of the theory behind and interpretation of pyroxene spectra in their investigation of the spectral effects resulting from pyroxene oxidation, as well as additional references. The following discussion has been summarized from the latter paper.

Pyroxenes have a structural formula of [(M1,M2)Si2O6. Metal cations occur in two crystallographic positions, the M1 and M2 sites. Pyroxenes in the Ludwig area are calcic clinopyroxenes of the diopside (CaMgSi2O6)-hedenbergite (CaFeSi2O6) series (Harris, 1979). In calcic clinopyroxenes, Fe2+ and Mg2+ occupy the M1 position and Ca2+ fills the M2 site. Spectra are dominated by Fe3+ absorptions near 0.95 and 1.15 μm. While these are weaker than absorptions caused by Fe3+ in the M2 site (e.g., pyroxenes of the enstatite-orthoferritesite series, (Fe, Mg, Si)O3), the greater crystal field stabilization energy imparted to the Fe2+ ion in clinopyroxenes results in absorptions that are considerably stronger than the Fe3+ bands present in garnets.

Bidirectional reflectance spectra of three skarn samples and four hornfels samples are presented in Figures 3a and 3b, respectively. The three skarn samples (YER3-5, YER3-33, and YER4-4) display progressively increasing garnet:pyroxene ratios. The hornfels samples (Figure 3b) are also arranged in order of decreasing pyroxene content from YER6-19 to MTE 502 and 514 to MTE 305.

**Figures 3.** Bidirectional reflectance spectra of garnet-pyroxene (a) skarn and (b) hornfels samples. Spectra have been offset for clarity; actual reflectance at 1.3 μm is labeled. Geoscan band centers are included.
Pyroxene-rich sample YER3-5 shows a strong, broad absorption centered near 1.1 μm, resulting from Fe²⁺ in pyroxene M1 sites. The reflectance maximum at 0.55 μm is not directly related to an electronic transition but is a "window" between two strong absorptions, the M1 crystal field at 1.1 μm and the intense Fe-O charge transfer in the UV. Sample YER3-33 (garnet-pyroxene ratio 60:40) still shows a weak pyroxene band at 1.1 μm, but the spectrum is now dominated by ferric absorption through 0.55 μm and at 0.87 μm. Sample YER4-4 is nearly pure andradite garnet, and the spectrum shows a well-developed 0.87 μm ferric band.

Sample YER6-19 (Figure 3b) is a pale green, fine-grained diopside hornfels with significant (20 to 30 percent) garnet. The spectrum exhibits two Fe²⁺ absorptions: a weak 0.95-μm and an asymmetric 1.1-μm feature. Sample MTE 514 consists of 80 to 90 percent isotropic garnet, with the remainder fine-grained pyroxene needles; the VNIR spectrum shows the 0.87-μm garnet absorption and a weak 1.2-μm band of indeterminate origin. Sample MTE 502 also consists of 80 to 90 percent isotropic garnet (0.2- to 1.0-mm diameter) with intimately intergrown pyroxene and epidote. The spectrum is nearly identical to that of MTE 514 but shows a slightly stronger 0.87-μm garnet absorption and a steeper slope toward the blue and UV. This is probably an effect of the protolith, a dolomitic marble. Hornfelses and skarn formed in pure carbonates tend to contain fewer precursor mineral inclusions and typically show stronger spectral features as a result. Sample MTE 305 formed in silty limestone and contains 80 to 85 percent anisotropic garnet with semi-opaque inclusions as the remainder. Only the garnet band at 0.87 μm is present in the VNIR.

The majority of the skarn and skarnoid in the study area are garnetites containing lesser, varying quantities (0 to 20 percent) of pyroxene. In spite of the much stronger spectral features of pyroxene, the spatial prevalence of garnet-rich rocks results in most skarn and hornfels spectra being dominated by the ferric iron garnet signature. In laboratory spectra measured from reconnaissance field samples, skarn tended to show a deeper 0.87-μm minimum than did hornfelses. This suggested that Geoscan channel 8 (0.873 μm) might be used to discriminate these alteration products.

In order to determine whether this difference in absorption depth was also present in the Geoscan data, pixel spectra were extracted from the image locations of these same seven samples (Figures 4a and 4b). These spectra are not strictly comparable with those in Figures 3a and 3b, as these laboratory spectra were measured from fresh surfaces and the image data represent weathered material. Nevertheless, the skarn samples in Figure 4a show a deeper local minimum in channel 8 than the hornfels samples in Figure 4b.

**Skarn/Hornfels Image Treatments and Results**

The treatment that proved most useful at differentiating between skarn and hornfels applied a (0.717-μm - 0.873-μm) band difference (Geoscan channels 5-8). The bright areas on the smoothed (5-8) single band image of the study area (Plate 2a) match well with the mapped distribution of skarn (Plate 2b). Only a few DN separate these anomalies from background, however; contrast stretching was used to identify areas that were obviously brighter than the norm.

A number of areas not mapped as skarn appear bright on this treatment and are annotated on Plate 2a. Several are small garnetite dumps from the Douglas Hill mine (1). Muscovite schists and pyritic hornfels in the andesite unit immediately east of the main skarn body (2) are heavily stained with iron oxides; limonite (goethite?) is the dominant mineral, but hematite is common and jarosite locally is present along fractures. The outcrop and dumps of the Ludwig silica-pyrite body (3) are similarly iron-stained, but hematite is less common and no jarosite was observed. These iron minerals show a strong crystal field absorption ranging from 0.87 μm (hematite) to 0.95 μm (goethite), just as do garnets containing ferric iron (Townsend, 1987). Strongly iron-stained rocks give a (5-8) value similar to that of garnet skarn, particularly if hematite is present.

**Possible Sources of Skarn/Hornfels Discrimination**

The observed spectral discrimination between the garnet-pyroxene skarn and garnet-pyroxene hornfelses could result from differences in composition, grain size, weathering styles, or any combination of the three. Garnets and pyroxenes in skarn are considerably more iron-rich than those in skarnoid; given that the (0.717-μm - 0.873-μm) treatment used above takes advantage of the 0.87-μm Fe²⁺ absorption in garnet, the higher (5-8) values in skarn may result from the higher iron content. It is well established that depth of an absorption feature increases with increasing grain size (Adams and Filice, 1967; Clark and Roush, 1984), a factor which would also favor a deeper absorption feature in the coarser-grained skarn. Weathering may affect skarn and hornfels differently and should be considered a critical factor in any study concerned with spectral differentiation.

Laboratory spectra were averaged over Geoscan band-passes for comparison with the airborne data. When analyzing laboratory spectra, ratios of reflectance are often
Compositional Effects

The 0.87-μm absorption feature results from Fe⁺⁺ within a mineral or in a weathering coat on the surface of a mineral. Given the dichotomy in iron content between Fe-rich skarn and Fe-poor hornfels, compositional variations must be con-
sidered as a possible source for the observed spectral discrimination. If the depth of the absorption feature increases with Fe content, it would fit the observed pattern of larger (5-8) values in skarn.

There is some empirical support for such a hypothesis. Moore and White (1972) studied the absorption spectra of nine garnets in the andradite-grossular-uvvarovite series as a function of their composition. They found a linear relationship between absorbance and weight percent Fe$_2$O$_3$ for five absorption bands between 0.35 and 0.75 μm. However, several factors prevent the direct application of their results to this study. Moore and White (1972) did not examine the 0.87-μm absorption band of interest in the Yerington garnets. They studied only shorter wavelength bands, none of which could be resolved in reflectance spectra of garnets used in this study. Their samples contained no garnets with intermediate compositions similar to those found in the Yerington hornfels; their grossularitic samples had compositions ranging from ad$_{20}$ to ad$_{26}$, while their andraditic samples had compositions of ad$_{35}$ and ad$_{45}$. Finally, they found absorbance to be a linear function of Fe$_2$O$_3$ content; reflectance, however, does not vary directly with absorbance and would not necessarily show a linear relationship. Nonetheless, their study lends credence to the hypothesis that the 0.87-μm absorption feature would deepen with increasing Fe.

To test this hypothesis, reflectance spectra averaged over Geoscan bandpasses were compared to composition for 16 garnet-pxyroxene skarn and hornfels samples in the study area. All samples consist of >70 percent garnet. Samples were selected that minimized spectral effects not related to garnet composition; all are free of opaque or inclusions and showed little or no alteration to epidote or other secondary minerals. Figure 5 shows the (5-8) values of convolved laboratory spectra versus the average molar percent Fe-endmember of garnets as determined by electron microprobe.

Figure 5. Plot of molar percent Fe-endmember composition of garnets vs. (5-8) values. The (5-8) values are calculated from laboratory spectra (0 to 100 percent) averaged over Geoscan bandpasses and multiplied by 2.55 to allow comparison with the 8-bit (0 to 255) Geoscan data. Bars represent one standard deviation. Open triangles represent hornfels, closed squares are skarn.

There is a definite trend toward an increasing (5-8) value of convolved lab spectra (DN) with Fe content, it would fit the observed pattern of larger (5-8) values in skarn.

Effects. If garnetite exhibited a tendency to fracture along rims of a given composition, a cut surface would not yield a spectrum representative of the "typical" garnetite hand sample. No such tendencies have been noted in this study area, but the possibility should not be disregarded in similar studies. More importantly, these are spectra of hand samples and the effects of grain size have not been removed. This factor is considered in the following section. While these caveats suggest that this relation may not be applicable to all skarn deposits, the data presented in Figure 5 indicate that the majority of the garnetiferous rocks at Yerington exhibit a trend toward stronger 0.87-μm absorption with increasing iron content.

Grain Size/Textural Effects

In a study of the reflectance characteristics of olivines as a function of composition, King and Ridley (1967) found systematic spectral variations with Fe/Mg ratios and with Ni impurities but cautioned:

"The area of an absorption band, i.e., the area under a straight-line continuum, is not simply related to the Fe/Mg ratio of the sample unless the grain size is severely restricted.... The area of the 1-μm absorption band is positively correlated with grain size fractions of a given olivine composition." (p. 11.468)

It is well established that the spectrum of a particulate material is strongly affected by its grain size (Adams and Filice, 1967; Clark and Roush, 1984). As grain size decreases, overall brightness increases and spectral features become less pronounced. The skarns in the study area definitely tend to be coarser grained than the skarnoid; an increase in the depth of the 0.87-μm feature with grain size could result in the larger (5-8) values for skarn.

Unfortunately, a comparison between grain size (estimated from thin section) and (5-8) value for 22 samples proved inconclusive. The data suggested that the depth of the 0.87-μm absorption increased with depth, but with a correlation coefficient of only 0.26. The strongest evidence for the importance of grain size effects on spectra lies in their identification in numerous other studies (Adams and Filice, 1967; Clark and Roush, 1984; Crowley, 1986; King and Ridley, 1987; Windeler and Lyon, 1991).

A common feature in the hornfelses is the occurrence of many fine-grained mineral inclusions in garnet; such samples were excluded from the comparison because the presence of numerous inclusions would make the estimation of the "average" grain size difficult. However, this is a textural effect that would effectively result in a finer grain size. Alternatively, mineral inclusions can be opaque materials that absorb across all wavelengths. Mineral inclusions in garnet are much more common in the hornfelses than in skarn and are yet another effect that would favor deeper spectral absorption features in skarn.

Weathering Effects

The rocks of the Yerington district are not deeply weathered. Outcrops are plentiful and most hilltops and hill-sides lacking outcrops are covered with a surficial layer of residual rock fragments (desert pavement) reflecting the underlying lithology.

Nonetheless, electromagnetic radiation in the visible and short wave infrared regions is most strongly affected by the top 50 μm of a surface and is thus highly sensitive to weathering (Buckingham and Somer, 1963). Figure 6 illustrates the importance of weathering on spectral features. The mean (5-8) values for laboratory spectra of skarn and hornfels samples averaged over Geoscan bandpasses were compared.
While the (5-8) values for skarn tend to be higher than those for hornfels, there is significant overlap. When the data are subdivided into spectra of fresh (broken surfaces, thin section heels) and weathered samples, however, the sets of weathered skarn and hornfels are distinctly more separable than the fresh.

In order to lessen possible sampling errors in this analysis, (5-8) values were compared for spectra of fresh and weathered surfaces of 28 samples (Figure 7). Weathering reduced the (5-8) value for all skarnoid samples examined, while for skarn samples weathering resulted in little or no change. In five samples (denoted by a dotted connecting line) weathering actually increased the (5-8) value in the skarn.

The skarn sample with the greatest drop in (5-8) value, YER7-44, is weakly coated with desert varnish and thus probably does not represent a typical fresh/weathered pair of spectra. The spectral differentiation between skarn and hornfels clearly is due in part to differences in their weathering characteristics.

The weathering characteristics of the garnet-pyroxene rocks in the study area vary with texture and mineralogy. As pyroxenes weather much more readily than garnets, their weathering properties would have a more immediate effect than those of the garnets. Fine-grained rocks develop surface coatings more readily than coarse-grained rocks. Garnet-pyroxene hornfelses in the study area typically weather to a brown or dark red-brown color suggestive of goethite. Pyroxene-rich skarn acquires a similar brown (goethitic) color of the fresh pyroxene. Garnet-dominated skarn shows fewer surface weathering effects than the finer-grained skarnoid; the weathered surfaces of garnet-pyroxene skarn darken slightly, while garnet skarn exhibits little color change and the glassy luster of the fresh garnets usually is preserved. Overall, the garnet-pyroxene hornfelses contain greater proportions of finer-grained pyroxene than skarn. The better-developed limonitic coats on skarnoid undoubtedly result from this abundance of readily-weathered, fine-grained pyroxenes.

Most calc-silicate altered rocks in the Ludwig area consist predominantly of garnet but contain significant pyroxene. For these garnet-rich rocks, spectra of weathered surfaces differ from fresh surfaces in the VNIR in several ways (Figures 8a and 8b):

1. The average albedo decreases;
2. The slope of the spectrum at wavelengths shorter than 0.7 μm may steepen; and
3. The Fe²⁺ absorption feature resulting from garnet is usually retained, but is often canted downward towards the shorter wavelengths as a result of (2). While this results in an apparent shift of the band location, isolation of the absorption feature by continuum removal proves this to be an illusory effect (Clark and Roush, 1984).

Whereas the garnet-rich rocks absorb strongly in the UV and thus also have relatively steep positive slopes into the visible, fine-grained iron oxides exhibit a steeper slope because of their “trans-opaque” behavior. At the longer wavelengths, reflectance increases with decreasing grain size (trans-), while at shorter wavelengths absorbance increases with decreasing grain size (opaque). The slope break between these two regions occurs near 0.55 μm (Salisbury and Hunt, 1968).

The spectra of pyroxene-rich rocks (Figure 8c) change similarly on weathering, although they differ from those of the weathered garnets due to differences in the original sample spectrum. Sample albedo decreases with weathering at wavelengths shorter than 0.6 μm but increases at longer wavelengths, again due to the trans-opaque behavior of the oxides. A study by Straub et al. (1991) determined that on oxidation of Fe²⁺ in calcic clinopyroxenes to Fe³⁺, the Fe³⁺ charge transfer transition (0.75 μm) masked the Fe²⁺ crystal field absorption feature. They also found evidence for nanophase (particle diameters <10 nm) hematite on oxidized surfaces. In this study, the strong, broad 0.95- and 1.1-μm Fe²⁺ absorption bands in the pyroxenes remain but are typically less well-defined than on fresh surfaces. An absorption may develop near 0.9 μm but is typically broad and shallow.

Effects (2) and (3) thus give a possible justification for the drop in (5-8) values between fresh and weathered hornfels samples illustrated in Figures 6 and 7. Fine-grained pyroxenes weather to iron oxides, and the spectral qualities of this oxide coat reduce the continuum reflectance in channel 5. The strongest effects of the limonite coat are seen in the visible region; channel 8 is less affected or unchanged by this coating and thus the (5-8) value decreases. As the inten-

![Figure 6. Average (5-8) values for convolved laboratory spectra of skarn and hornfels, fresh and weathered. Horizontal ticks are means, vertical bars represent one standard deviation.](image)

![Figure 7. Pairs of (5-8) values for fresh and weathered surfaces of 12 skarn and 16 hornfels samples. Open symbols represent fresh surfaces, closed are weathered. Solid connecting lines indicate a decrease in (5-8) on weathering, dotted lines indicate an increase.](image)
Conclusions
Thermal IR difference images were successful in defining the distribution of garnet-pyroxene skarn and hornfels in the Ludwig study area, while a (0.717-μm - 0.873-μm; 5-8) difference image provided a means of differentiating between the two. Areas heavily stained by iron oxides were present as errors of inclusion on the (5-8) image, but the additional information gained through the thermal IR differences could readily be included to distinguish these from skarn. One simple method would be a (23-20) [5-8] [23-19] image (Winderler, 1992). Alternatively, this combination of thermal and VNIR band differences could be used as a means of identifying iron-stained, silicified rocks - common targets in mineral exploration.

Compositional, grain size, and weathering variations were investigated as possible sources for the skarn/hornfels discrimination displayed by the (0.717-μm - 0.873-μm; 5-8) difference image. All three factors contribute, but spectral variations due to weathering appear to be the most important. These results are summarized in Table 4.

Skarn/hornfels discrimination in the Yerington contact aureole thus appears to result from several spectral variations that fortuitously all contribute in the same direction. Whether this treatment would be effective in other districts is uncertain, although higher iron contents and coarser grain size relative to early hornfels are common features of skarn districts (Einaudi et al., 1981). The VNIR spectral effects of manganese may also play a role, particularly in Pb-Zn skarns. However, the use of multispectral thermal infrared data for detection of garnetiferous rocks should be generally applicable to skarn exploration. Most skarns are associated with felsic to intermediate intrusions, which would have emission spectra distinct from the relatively long-wavelength minima of garnetites in the 8- to 13-μm region. Discrimination between different stages of calc-silicate alteration may often depend on site-specific, empirically determined treatments.

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References
<table>
<thead>
<tr>
<th>Possible source of discrimination</th>
<th>Theorized effect on spectra</th>
<th>Why considered in this study</th>
<th>Actual effect on spectra</th>
</tr>
</thead>
<tbody>
<tr>
<td>Iron content</td>
<td>Minerals with greater Fe-content may show stronger Fe-absorption</td>
<td>Skarn garnets are considerably more iron-rich than hornfels</td>
<td>Some effect; garnets show increase in depth of Fe$^{2+}$ minimum at 0.873 μm</td>
</tr>
<tr>
<td>Grain size</td>
<td>Coarser grain size results in deeper absorption minima</td>
<td>Skarn is generally coarser-grained than hornfels</td>
<td>Probably contributes, but by itself is only a minor factor</td>
</tr>
<tr>
<td>Weathering styles</td>
<td>Different weathering styles or products may result in different spectra</td>
<td>Skarn and hornfels contain different grain size and quantities of garnet &amp; pyroxene</td>
<td>Major factor in discrimination; greater quantity of fine-grained pyroxenes in hornfels weather easily and reduce brightness in channel 5 relative to channel 8</td>
</tr>
</tbody>
</table>


