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#### Notes, Comments, and New Findings

#### Semantic Categorization and High-Speed Scanning

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Two experiments used versions of Sternberg's item-recognition task, in which the subject could make a decision by semantic categorization or by a search of short-term memory. When a single category distinguished the words of a memory set from foils, pronounced deviations from a linear set-size effect were observed. Decision times were affected little by increases in set size beyond three or four, suggesting that a categorization process was circumventing an item-by-item search of memory. These results were observed in the absence of a consistent mapping between stimuli and responses; in other words, within a session the relevant category constantly changed and stimulus words were used as both targets and foils. The results are compatible with a race model in which memory scanning and semantic categorization occur in parallel; the decision time is determined by the process reaching completion first.

Sternberg (1966) reported that a decision based on a search of memory produced decision times that increased linearly with the number of items to be searched. For small memory loads, this setsize effect has proven remarkably robust across a variety of item types, including letters (Nickerson, 1966), digits (Sternberg, 1966), words (Smith, 1967), and random forms (Sternberg, 1969). Sternberg concluded that subjects serially scan through the items of short-term memory (STM).

Experiments demonstrating this linear set-size effect use an item-recognition paradigm. In this paradigm, the subject commits a small set of items (usually six or fewer) to memory and must then make a speeded yes/no response regarding the presence or absence of a test item in this set. Targets and foils are commonly all of the same item type, and no feature reliably distinguishes them. In the absence of such a feature, subjects are forced to base their decision on a search of STM.

In experiments in which one or more features do distinguish targets and foils, there is often a substantially reduced and sometimes curvilinear relationship between decision time and set size. This effect was observed by Ellis and Chase (1971) at the perceptual level when foil letters were distinguished from targets by the features of size or color. A letter/digit categorical distinction has been used to similar advantage in the item-recognition task (Lively & Sanford, 1972; Simpson, 1972). In an extension of the item-recognition task, Schneider and Shiffrin (1977) observed almost no effect for set size when targets and foils were separated by the letter/digit distinction. Semantic categorical membership can also apparently facilitate the rejection of foil words (Okada & Burrows, 1973; Reynolds & Goldstein, 1974).

These experiments suggest that decisions are sometimes made on the basis of a categorization process that is relatively unaffected by set size. Such a process would be increasingly likely to be completed before an item-by-item search as set size increases, resulting in the often observed curvilinear relationship between decision time and set size. However, the design in each of the abovementioned experiments creates a form of response consistency. In all of these experiments, for example, a proportion of the foils are drawn from a categorically defined set and are never used as targets. Schneider and Shiffrin (1977) argued that response consistency permits the development of an automatic detection response that is unaffected by set size. The development of automatic detection is speeded by a target/foil distinction but does not depend on it.

The present study investigated the effects of semantic target/foil distinctions in the absence of response consistency. A central question con-

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Table 1 Stimulus Words for Experiments 1 and 2

HOUR SECOND YEAR	HORSE LION RABBIT	BLUE GREEN YELLOW	HEAD FINGER NOSE	SHIRT PANTS	PIANO DRUM
MINUTE	DEER	PURPLE	TOOTH	BLOUSE COAT	VIOLIN FLUTE
DECADE WEEK	SHEEP BEAR	WHITE PINK	KNEE ANKLE	DRESS JACKET	GUITAR TUBA

cerned the extent to which a categorization process could replace the scanning process as a basis for decisions. Given the speeds with which scanning can take place (much less than I see for sets of fewer than six words), such a replacement would demonstrate a very rapid encoding and utilization of semantic information.

#### Experiment 1

#### Method

Materials and design. Memory sets, targets, and foils were all drawn from a stimulus set consisting of six exemplars from each of six categories: units of time, four-legged animals, colors, parts of the human body, articles of clothing, and musical instruments (see Table 1). The words were the most common members of their respective categories, as determined by the Battig and Montague (1969) word norms with the following restrictions: (a) Each word was three to eight letters in length. (b) No more than two words of a category began with the same letter. (c) No word belonged to more than one category of the Battig and Montague norms (e.g., orange). (d) No pairs of words were graphemically or phonemically confusable (e.g., mouth and mouse).

A usable target/foil distinction was established through a manipulation of memory-set type. For sets of multicategory set type, each item represented a different category; in single-category sets all items came from the same category. The factors of set size, test period, and response type were crossed with the factor of set type in a withinsubjects design. The size of a memory set ranged from one to six words; an experimental session consisted of six test periods. Within a test period, each of 12 memory sets was tested in a block of 6 targets and 6 foils. The 12 memory sets represented each combination of set size by set type. Each single-category set was randomly selected such that no two sets of a test period came from the same category. Multicategory sets were similarly selected so that no group of words occurred in more than one set.

Within a test block, each of three foils was pre-

sented twice. The number of presentations for a memory-set item was in proportion to the set size (approximately, six divided by the set size). Each foil in a test block came from a different category; for single-category sets, none of these categories matched the category of the memory set. Each stimulus word had an equal chance of appearing in any combination of set type by response type within a test period. In an experimental session a word's use as a foil always roughly equalled its use as a target.

Apparatus. The experiment was run on a PDP 11/34 computer using the RSX-11M system. All stimuli were displayed in uppercase letters on a Beehive 100 terminal using a 5 × 7 (per character) dot matrix. The terminal was modified to display stimuli only at the beginning of a video frame; all stimulus-dependent timing was initiated at the beginning of a frame. Yes/no responses were made through a hand-held, two-button box. The left button was labeled "no" and the right button was labeled "yes."

Subjects and procedure. Seven male and six female subjects between the ages of 18 and 26 participated in a single 2-hr. session and were paid between \$5 and \$8 for their participation. A game format was used in which subjects were awarded points for fast errorless performance; these points were converted into money at the end of the experimental session. Subjects earned a half point for each decision time faster than a "time to beat" (discussed later), and subjects lost two points for each error.

The subjects became familiar with the words and categories through three initial tasks. In an old/new recognition task, each word of the stimulus array was presented twice in a random order and subjects made the appropriate speeded response. In a spelling task, each word was presented for 1 sec, after which the subjects were required to correctly spell the word. A spelling mistake initiated an immediate re-presentation of the word. In an STM span task, the six exemplars of a category were presented, in succession, at 1-sec intervals, and the subject attempted to recall the words in any order. Less than perfect recall brought about an immediate re-presentation of

the exemplars in a new order. Each categoriested to a criterion of one perfect recall the examplars of the category were correcalled in this manner the subjects were protype in a name for the category.

The instructions preceding the item-rece task (a) informed subjects that a categor: tinction between targets and foils would times exist, and (b) encouraged them to m of this distinction. Two practice test blotiated each test period. These blocks gave ject the opportunity to win bonus points, \*\* subject's average decision time for these determined the initial time to beat. Before of the remaining test blocks in a period, t to beat was set equal to the smaller of tw ers: (a) the previous time to beat multi; 05 msec, or (b) the average of the previo beat and the actual mean decision time revious block. This number was then a r variations in set size by adding the fac $ze - 3) \times 25$  msec.

Each test block began with a screen dia) the memory set presented in a rando dered column, and (b) the time to beat. Fiewed the display for as long as they wi press of the "return" button on the termi board terminated the display and initiate omly ordered presentation of the test iter ding the display of each test item, the ADY appeared on the screen, followed b ec delay. The screen then cleared, a to peared, and the timer began. A press of no response button terminated the dist topped the timer. At the end of a test blo ets were informed of their average decis: be number of times they were faster t me to beat, their error rate, their points and their new point total.

Within a test period, memory sets wated randomly with respect to set size pe. After every other test period, subjected a span task and were then given teak. The span task's primary purposeak the tedium of the item-recognition tean-task results are not reported here.

#### esuit.

Decision times and error rates were so an analysis of variance. Decision timecarded for trials in which an error we the time itself exceeded I sec. Less the correct responses were longer that the time I shows the relationship between the and set size for the four combinative and response type.

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the exemplars in a new order. Each category was tested to a criterion of one perfect recall. After the examplars of the category were correctly recalled in this manner the subjects were prompted to type in a name for the category.

The instructions preceding the item-recognition task (a) informed subjects that a categorical distinction between targets and foils would sometimes exist, and (b) encouraged them to make use of this distinction. Two practice test blocks initiated each test period. These blocks gave a subject the opportunity to win bonus points, and the subject's average decision time for these blocks determined the initial time to beat. Before each of the remaining test blocks in a period, the time to beat was set equal to the smaller of two numbers: (a) the previous time to beat multiplied by 105 msec, or (b) the average of the previous time to beat and the actual mean decision time for the previous block. This number was then adjusted for variations in set size by adding the factor (set  $_{\rm size}$  - 3)  $\times$  25 msec.

Each test block began with a screen display of (a) the memory set presented in a randomly ordered column, and (b) the time to beat. Subjects viewed the display for as long as they wished. A press of the "return" button on the terminal keyboard terminated the display and initiated a randomly ordered presentation of the test items. Preceding the display of each test item, the word READY appeared on the screen, followed by a 500msec delay. The screen then cleared, a test item appeared, and the timer began. A press of the ves or no response button terminated the display and stopped the timer. At the end of a test block, subects were informed of their average decision time, the number of times they were faster than the time to beat, their error rate, their points earned, and their new point total.

Within a test period, memory sets were presented randomly with respect to set size and set type. After every other test period, subjects performed a span task and were then given a 3-min. break. The span task's primary purpose was to break the tedium of the item-recognition task, and span-task results are not reported here.

#### Results

Decision times and error rates were submitted to an analysis of variance. Decision times were discarded for trials in which an error was made or the time itself exceeded 1 sec. Less than 1% of the correct responses were longer than I sec. Figure 1 shows the relationship between performance and set size for the four combinations of set type and response type.

With respect to decision time, main effects in-

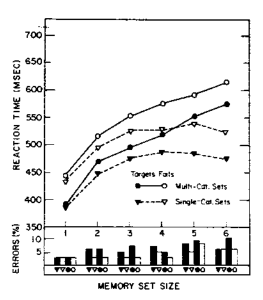


Figure 1. Data from Experiment 1: mean decision times for correct responses and percentages of error as functions of memory-set size (collapsing across test periods).

dicating the advantages of single-category sets, targets, and practice (across test periods) were all significant (p < .01). In addition, there was a setsize effect, F(5, 60) = 114.00,  $\rho < .001$ . However, there was a marked interaction between set size and set type, F(5, 60) = 22.63, p < .001. The linear component to the relationship between set size and decision time accounts for 92.9% of the variance in the multicategory condition but for only 63.1% of the variance in the single-category condition. In contrast, the quadratic component accounts for 5.6% of the variance in the multicategory condition and for 35.8% of the variance in the single-category condition.

The departure from linearity for the single-category condition can be attributed to an apparent lack of any effect when set size was increased beyond three words. The mean time for a correct yes response in the single-category condition was 473 msec for three-item sets and 475 msec for sixitem sets. The mean times for a correct no response were 525 msec and 521 msec for these same two set sizes, respectively.

In the accuracy data, the main effects of set size and response type were both significant (p <.02). In agreement with the decision-time results, there was a marginally significant interaction between set size and set type, F(5, 60) = 1.95, p < .10. There is no indication, then, that the decision-time results can be explained as a simple speed/accuracy trade-off. The overall error rate

Table 2
Stimulus Words Used to Extend the Stimulus Set for Experiment 2

MONTH	TIGER	DDOWN			
	HOER	BROWN	FOOT	SOCKS	HARP
DAY	MULE	GRAY	LEG\$	BELT	TRUMPET
CENTURY	MOUSE	BLACK	ARMS	HAT	BANJO

#### Discussion

A categorical distinction, in the absence of response consistency, has a clear-cut and dramatic effect on performance in the item-recognition task. The curvilinear relationship between decision time and set size for single-category sets strongly implies a categorization process, which is unaffected by set size.

Ellis and Chase (1971) proposed that categorization at a perceptual level and scanning could take place in parallel, with decision times reflecting the faster to complete of the two processes. In tasks requiring a dual judgment for each item of the memory set, Burrows and Okada (1973, 1976) and Palef (1977) found evidence that semantic information can also be retrieved in parallel with a serial scan.

Categorization and scanning, however, need not take place in parallel to account for the data. Another possible model uses a probabilistic mixture of categorization and scanning (e.g., Atkinson, Herrmann, & Wescourt, 1974). In such a model, the probability that categorization is first attempted for single-category sets of three or more items would have to be near 100% to account for the present results. In fact, subjects in the experiment could have made a strategic commitment to categorization at the beginning of a single-category test block. If they chose to make their decisions on a categorical basis, there would have been no need to even maintain the individual set items in memory, since categorization alone would always have provided the correct response.

The second experiment made such an a priori commitment to categorization impossible. The experiment tested single-category sets in a test block containing an equal number of related foils (same category) and unrelated foils (different categories). Since presentation order for the two foil types was random, the subject would always have to keep the memory set active in order to distinguish targets from related foils. However, it would be possible to reject unrelated foils by means of a category judgment. Of particular interest is the interaction between scanning and categorization in a task where the majority of test items in a block require a serial scan.

#### Experiment 2

#### Method

Materials and design. All items of a memory set were drawn from the same semantic category. The factor of foil relatedness replaced the set-type manipulation of Experiment 1. Three foils in a test block were categorically related to the items of the memory set, and the remaining three foils were unrelated. The use of related foils with larger memory sets required an increase in the number of exemplars per category; this was accomplished by adding the words shown in Table 2.

As in the previous experiment, there were six test periods with 12 blocks each. Within a test period, two memory sets of different size were randomly selected from each of the six categories. Three related foils were randomly drawn from the words remaining in a category after a set's selection. The three unrelated foils were randomly drawn from a random collection of three of the five remaining categories. Within an experimental session, each word had an equal chance of appearing as a target and a foil. In addition, each word had an equal chance of appearing as a related and unrelated foil.

Subjects and procedure. Seven male and eight female subjects from the same population used in Experiment I participated in a single 2-hr. session and were paid between \$5 and \$8 for their participation. The familiarization tasks again preceded the item-recognition task. In the span-familiarization task, the word sequence length was reduced from six to five, and each category in Table 2 was tested twice to insure that each word was included in at least one span sequence.

In the instructions preceding the item-recognition task, subjects were encouraged to use category information in the rejection of unrelated foils. In all other respects, experimental procedure and design followed that of Experiment 1.

#### Result

Figure 2 shows the relationship between performance and set size for the three test-item types (targets, related foils, and unrelated foils). Decision times were screened using the procedure described for Experiment 1. Again, fewer than 1%

of the correct responses exceeded 1 sec. T for three subjects were excluded from the a because of an error rate exceeding 10%, arate analysis of the data for these three produced complete agreement with the from the remaining subjects, which follow

An analysis of the decision times revernificant main effects of practice (across riods) and set size (p < .001). Unrelated fo rejected significantly faster than relate were, F(1, 11) = 130.43, p < .001. The diff effect of set size on the three test-item reflected in a significant interaction, F(10)22.97, p < .001. In order to understand teraction, the two foil conditions were set considered in relation to targets. The effect was significantly less for unrelated fo was for targets, F(5, 55) = 16.95, p < 0the other hand, the effect of set size was antly greater for related foils than it was lets, F(5, 55) = 3.57, p < .01. The line: onent of the relationship between set s ecision time accounted for 98.9% of the v or related foils and 91.7% of the varia nrelated foils. The quadratic compon counted for less than 1% of the variance iated foils and for 7.9% of the variance related foils. For targets, the linear conecounted for 92.3% of the variance and t atic component accounted for 6.1% of t ce. These figures for targets are in close ent with the corresponding figures t palticategory condition of Experiment 1 In the error data, a main effect for set : mificant, F(5, 55) = 15.13, p < .001. sion-time advantage for unrelated over oils was reflected in error rates as we (1) = 13.24, p < .004. Set size had a signi reater impact on error rates for related f or targets, F(5, 55) = 5.48, p < .001. The significant interaction in the relation een set size and targets versus unrela-> .17). However, error rates for targe gnificantly greater than for unrelated fo (1) = 46.75, p < .001. The error rates for the following the first the lowest of the following paditions and were obviously unaffected cases in set size. There is no indicat cision-time differences are solely the 1 speed/accuracy trade-off.

#### scussion

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#### Experiment 2

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2 shows the relationship between perand set size for the three test-item types related foils, and unrelated foils). Dees were screened using the procedure for Experiment 1. Again, fewer than 1% of the correct responses exceeded 1 sec. The data for three subjects were excluded from the analysis because of an error rate exceeding 10%. (A separate analysis of the data for these three subjects produced complete agreement with the results from the remaining subjects, which follow.)

An analysis of the decision times revealed significant main effects of practice (across test periods) and set size (p < .001). Unrelated foils were rejected significantly faster than related foils were, F(1, 11) = 130.43, p < .001. The differential effect of set size on the three test-item types is reflected in a significant interaction, F(10, 110) =22.97, p < .001. In order to understand this interaction, the two foil conditions were separately considered in relation to targets. The effect of set size was significantly less for unrelated foils than it was for targets, F(5, 55) = 16.95, p < .001. On the other hand, the effect of set size was significantly greater for related foils than it was for targets, F(5, 55) = 3.57, p < .01. The linear component of the relationship between set size and decision time accounted for 98.9% of the variance for related foils and 91.7% of the variance for unrelated foils. The quadratic compenent accounted for less than 1% of the variance for related foils and for 7.9% of the variance for unrelated foils. For targets, the linear component accounted for 92.3% of the variance and the quadratic component accounted for 6.1% of the variance. These figures for targets are in close agreement with the corresponding figures for the multicategory condition of Experiment 1.

In the error data, a main effect for set size was significant, F(5, 55) = 15.13, p < .001. The decision-time advantage for unrelated over related foils was reflected in error rates as well, F(1,11) = 13.24, p < .004. Set size had a significantly greater impact on error rates for related foils than for targets, F(5, 55) = 5.48, p < .001. There was no significant interaction in the relationship between set size and targets versus unrelated foils (p > .17). However, error rates for targets were significantly greater than for unrelated foils, F(1,11) = 46.75, p < .001. The error rates for unrelated foils were by far the lowest of the three conditions and were obviously unaffected by increases in set size. There is no indication that decision-time differences are solely the result of a speed/accuracy trade-off.

#### Discussion

Although most of the trials in a test block required a memory scan, the categorization process continued to facilitate the rejection of unrelated foils, especially at the larger set sizes. The total increase in decision time as set size ranged from

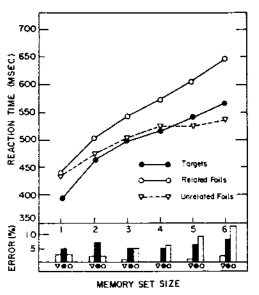


Figure 2. Data from Experiment 2: mean decision times for correct responses and percentages of error as functions of memory-set size (collapsing across test periods).

four to six was only 15 msec for unrelated foils, in comparison to a 74-msec increase for related foils and a 44-msec increase for targets. The average time to reject an unrelated foil at these larger set sizes is virtually identical to the rejection time for single-category foils in Experiment 1 (528 vs. 529 msec). The greater linearity of the decision-time slope may indicate that decisions for unrelated foils were sometimes based on a serial scan for even the larger set sizes.

Both the decision-time and error-rate slopes are significantly steeper for related foils than for targets, suggesting that the category membership of a foil may impede as well as facilitate its rejection. This is further indicated by a comparison of Experiments 1 and 2, in which differences in performance are noticeable for sets of five and six items. For these set sizes, targets in Experiment 2 are about 12 msec faster than multicategory targets in Experiment 1. In contrast, related foils in Experiment 2 are about 24 msec slower than multicategory foils in Experiment 1. In addition,

If the decision-time slope for related foils is depressed as a result of a speed/accuracy trade-off (a possibility suggested by the high error rates for related foils), then the slope may actually be in a 2:1 relationship with the decision-time slope for targets. Therefore, subjects may have used a terminating search to distinguish targets from related foils. However, it is not clear why a terminating search would selectively produce such high error rates for related foils.

error rates are considerably higher for related foils at the larger set sizes. Poor performance for related foils could occur in a parallel model as the result of a response competition between the categorization process and scanning. It is likely that a suppression of the category yes response would take extra time; a failure to suppress this response would lead to an error.

#### General Discussion

The results of these two experiments are perhaps best described in terms of a parallel race model involving serial scanning and the categorization process. Categorization dominated for sets of more than three or four items and was unaffected by changes in set size. These effects were observed in the absence of a consistent mapping between stimuli and responses. In both experiments the relevant category changed after every 12 trials, and stimulus words were used equally often as targets and foils.

The results demonstrate a use of semantic information to circumvent an item-by-item search of memory. Similar results have been obtained in the fact-retrieval paradigm of long-term memory (McCloskey & Bigler, 1980; Reder & Anderson, 1980; Smith, Adams, & Schorr, 1978). In the experiments of Reder and Anderson, for example, interference did not occur among a set of thematically related facts when foils were thematically unrelated. Interference returned, however, when thematically related foils were used.

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Strategie

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Adult subjects' strategies in twere studied in three experim of either an array rotating relative to a fixed ar logically equivalent problems lems differed in overall difficuterns. The data suggested that problems. For viewer rotation a more abstract frame of refersonly one element at a time-functional "unit" of spatial tr

In the past decade there has been a treigrowth of interest in the processes involve tial reasoning. Much of this interest has a from studies of the "mental rotation" ponon by Shepard and his colleagues (e.g. & Shepard, 1975; Shepard & Metzler, 19 variety of related tasks, chronometric desupported a powerful process model of homotopy manipulate spatial informations have generally dealt with figures per chistoscopically, and the mental rotatic roposes that adults holistically rotate a res in an analog fashion.

In the current experiments, adult spanning was examined across a wider rangemands than has typically been studingically equivalent spatial reasoning perfect compared using object arrays and questioning procedures. In the first type m (directly akin to those studied by Subjects predicted the outcome of a rotat tray relative to a stationary viewer (ation problem). In the second problem, redicted the outcome of a rotation of tound the periphery of a stationary array otation problem).

These array-rotation and viewer-rotatems were contrasted for three types of g procedure that probed different aspearsformed spatial information. The three types were termed appearance, item

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# Interaction between water and melt in the system $CaAl_2O_4$ -SiO<sub>2</sub>- $H_2O$

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(Received November 7, 1989; revised and necepted May 15, 1990)

#### BSTRACT

Mysen, B.O., 1990. Interaction between water and melt in the system ([aAl<sub>2</sub>O<sub>4</sub>-SiO<sub>2</sub>-H<sub>2</sub>O<sub>2</sub>Chem. Geol., 88: 223-243)

The interaction between dissolved  $H_2O$  and such structure on the join  $(\pi A|_2O_4 - SiO_2 + H_2O)$  has been studied with Raman spectroscopy. The total  $H_2O$  contents ranged from 3 to 10 wd.% with AI/(AI + Si) = 0-0.33. The spectra are consistent with formation of OH complexes that include all  $(\pi^2)$  and  $AI/(\pi)$  in addition to molecular  $H_2O$ . No direct evidence for  $(S_1A) + OH$  bonds can be discerned in the spectra of  $(\pi^2) + OH$  stretching observed in the spectra of  $(S_1A) + OH$  direct in aluminous samples). However, the spectral topology of the fundamental OH stretch bands near 3600 cm. Can only be rationalized if some Si+OH or  $(S_1A) + OH$  bonding exists in the metts.

The melts become depolymerized as H<sub>2</sub>O is dissolved to form Ca OH and ALOH complexes. Formation of Ca.OH complexes is a more efficient depolymerization mechanism than that of ALOH complexes [6 vs. } nonbridging oxygen would be formed per mole H<sub>2</sub>O dissolved as a Ca.OH complex of Ca(OH), type vs. an ALOH complex of A{OH}, type W. an ALOH complex of A{OH}, type W. an ALOH complex of A{OH}, type W. an ALOH complex of Definition of the melts complexing of OH with AP<sup>2</sup> (ALOH) probably becomes more important at the expense of complexes with Ca<sup>2</sup> (Ca. OH). Thus, the effect of dissolved H<sub>2</sub>O on melt polymerization diminishes with Al/(Al+Si). However, the degree of polymerization of the melts (NBO/T) for a given total H<sub>2</sub>O concentration is less than that expected by either the Ca.OH or the ALOH complexing nucchanism alone. The excess water is present as molecular H<sub>2</sub>O and as (Si,Al)-OH bonds that replace (Si,Al)-OH(Si,Al) bridging oxygen bonds in the melts.

#### i. introduction

Dissolved water affects chemical and physial properties of silicate metts. For example, transport properties are strongly enhanced in hydrous melts compared with their anhydrous quivalents. Melt viscosity can be lowered by upto several orders of magnitude (e.g., Shaw, 1963; Kushiro, 1978; Dingwell and Mysen, 1985). Cation diffusivity is increased (Walson, 1979) as is electrical conductivity (Takatatal., 1981; Satherley and Smedley, 1985). Uquidus relations are also significantly affected by water, generally resulting in a decrease in silica activity (e.g., Kushiro et al., 1988; Kushiro, 1972, 1989; Mysen and Boettcher, 1975; Carmichael et al., 1976).

OH bonding types in silicate melts have been and Oxtoby, 1986) are suggestive of multiple tems (Oxtoby and Hamilton, 1978; Hamilton with OH groups and an exchange of H+ from tion, molecular H<sub>2</sub>O can be important in sili inferred (Mysen and Virgo, 1986a). In addi-Na<sub>2</sub>O-Al<sub>2</sub>O<sub>3</sub>-SiO<sub>2</sub> system, a variety of metal-OH species. From spectroscopic studies in the solubility data for alkali aluminosilicate sys-H<sub>2</sub>O with Na<sup>+</sup> in the melt (Uys and King, through the replacement of bridging oxygen melt have have led to inference that the melts surements of hydrous NaAlSi<sub>3</sub>O<sub>8</sub> composition 1963; Burnham, 1974, 1975). However, water become depolymerized upon solution of  ${
m H_2O}$ and water activity together with P-V-T mca-The relationships between melt properties

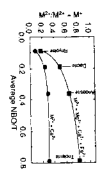


Fig. 1. Proportion of divalent metal cations (M2+) relalation of NBO/T as described by Mysen (1988). Chayes (1975a, b. and unpublished data, 1985). Calcugree of polymerization (NBO/T). Chemical analyses from mon natural magmatic liquids as a function of their detive to monovalent and divalent metal cations in com-

cussion (e.g., Dingwell and Webb, 1989a, b; and Burnham, 1984; Epel'baum, 1985; Naal., 1975; Persikov, 1975; Stolper, 1982; Eggler cate melts (e.g., Hodges, 1974; Epel'baum et Silver and Stolper, 1989). H<sub>2</sub>O and OH groups remains a matter of dis-However, the relative abundance of molecular vrotsky, 1987; Silver and Stolper, 1989).

action between water and natural magmatic cations are represented by Ca2+ (Mysen common magmatic liquids are divalent (Ca<sup>2+</sup>, Mg<sup>2+</sup> and Fe<sup>2+</sup>). Over 50% of the divalent tion of rhyolite,  $\geq 50\%$  of the metal cations in in magmatic liquids (Fig. 1). With the excep-Na ' in particular, are not the principal cations atkali metals in general (K+ and Na+) and of the role of water in magmatic liquids, but join NaAlO2-SiO2, Data in this system are were alkali aluminosilicates, commonly on the in silicate melts is that the base compositions on solubility and solubility mechanisms of H2O this direction, interactions between H<sub>2</sub>O and aluminosilicate melts are needed. As a step in evident that in order to understand the inter-1988). From these simple observations, it is fundamentally important for characterization melts in the system CaO-Al<sub>2</sub>O<sub>3</sub>-SiO<sub>2</sub> have liquids, experimental data for alkaline-carth A common feature of the majority of studies

# 2. Experimental methods

Starting materials were glasses formed by

carbonation of CaCO3 at ~1300°C in vertical water was double-distilled and deionized beal. (1986) and Mysen and Virgo (1986a). The commercial sitica glass also used by Hemley et 0.200, 0.25 and 0.333 for the compositions CA2S8, CA2S6 and CA2S4, are on the join et al. (1982). All compositions, CA2S14 quench furnaces. Detailed chemical character at 1550-1660°C for 1-4 hr. subsequent to de melting CaCO<sub>3</sub>+Al<sub>2</sub>O<sub>3</sub>+SiO<sub>2</sub> oxide mixture contents ranged from 3 to 10 wt.%, which are water concentration in the sample from the tively. The SiO2 starting material was the same CA2S14, CA2S8, CA2S6 and CA2S4, respec- $CaAl_2O_4$ -SiO<sub>2</sub> with their Al/(Al+Si)=0.125 ization of these materials were given in Seifer ature conditions of synthesis (McMillan et al. compositions under the pressure and temperless than the water solubility in these melt combined weighting errors is  $\leq \pm$  5%. The H<sub>2</sub>O ±0.05 µl precision. The uncertainty of the into Pt capsules with a microsyringe with a mainder as light water. The water was loaded fore use, and D<sub>2</sub>O was 95.7% D<sub>2</sub>O with the re-1986).

gether with known amount of H2O (or D2O) and England, 1960). About 20-mg samples tosolid-media, high-pressure apparatus (Boyd ranging from 1.5 to 64 hr. (Table 1) in 0.5-in. kbar and 1500-1700°C with run duration in sealed Pt capsules were subjected to 13-15 s<sup>-1</sup> rate is shown as dashed lines in Fig. 2). distinct nonlinear functions of time since ditions with the 0.5-in. (1.27 cm) diameter power to the furnace) from experimental confurnace assemblies without external control of (1.27 cm) diameter Pyrex® glass-AlSiMag near 100°C s. I from the experimental temin. (17,78 cm) diameter by 2-in. (5.08 cm) furnace assemblies in steel pressure plates of 7the  $f_{\rm H_2}$ . The quenching rates (by turning off the peratures between 500° and  $600^{\circ}\mathrm{C}$  (the  $100^{\circ}\mathrm{C}$ quench (Fig. 2). In general, though, the rate is thickness (water-cooled above and below) are The hydrous glasses were produced in the

# INTERACTION RELIWEEN WATER AND AIL LINE GALJO, SIOT HJO

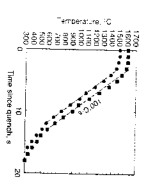
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Synabe	is condition	Synthesis conditions of high-pressure, hydrous glasses	essure, hyd	rous glasse	×
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E E	Starting	Volatile	Pressure	Pressure Temper- Run	Z III
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sperimental conditions (McMillan et al., ontent of the experimental samples) under the nelts exceeds 10 wt.% (the maximum H<sub>2</sub>O wenching because the water solubility in these when observed) are likely to be formed upon non of water upon quenching. The bubbles ated bubbles that most likely reflect exsoluchibited a tendency to form finely dissemicopic scale) glass. Samples with 10 wt.% 11.0 m clear, bubble-free (on an optical micro-All but the most water-rich samples quench

Structural information was obtained with an



quench) of solid-media, high-pressure apparatus used in Fig. 2. Quenching rates (expressed as temperature since the experiments,

Wat the ~1-mm' samples. both cases, the samples were excited with  $\sim 0.5$ line of an Ar' laser for sample excitation. In quency-doubled Nd-YAG laser and the 480-nm sen et al., 1982) with the 532-nm line of a freautomated Raman spectroscopy system (My

deuterated samples, the latter region is 2000cludes the range of OH stretch vibrations. For cm 1) includes the range of (Si,Al)-O and 3000 cm 1 frequency segment (3000-4000 cm. ') inregion of possible H-O-H bending. The higher-(Si,A1)=OH vibrations, and the 1600-cm<sup>-1</sup> The lower-frequency segment (300-1800 The spectra were recorded in two segments

the method of minimization of least squares et al., 1982) was carried out statistically with curve-fitting (Seifert et al., 1981, 1982; Mysen point of the greatest absolute intensity. The fitted spectrum were normalized to the data quency-dependent scattering intensity (e.g., were corrected for temperature- and freserved. Prior to statistical analysis, the spectra quencies where Raman scattering was ob-Long, 1977). The Raman intensities in each et al., 1982). The background was subtracted spectra were curve-fitted statistically (Mysen fitting of a line through the data points at frefrom the uncorrected spectra by least-squares tural information that could be obtained, the In order to maximize the amount of struc

Upon convergence, the minimum value of  $\chi^2$ quency, half-width and intensity) as well as the bution are obtained. All line parameters (freand maximum randomness in residual distrithe fitting routine. number of lines were independent variables in (Fletcher and Powell, 1963; Davidson, 1966).

convergence is global or regional; and (3) desirable to ascertain the reliability of the fita successful comparison of structural inforwas concluded that the results of the fitting whether the number of lines in the final fit is, chosen line-shape is appropriate; (2) whether, necessary to address (1) whether or not the are not always visually evident. Therefore, it is netic resonance (NMR) spectroscopy added mation with data obtained by nuclear magroutine fulfill these requirements. In that study, minosilicate systems (Mysen, 1990), where it anhydrous, depolymerized melts in alkali alu- $\chi^2$  and maximum randomness in the residuals. in fact the number that leads to the minimum for a given fit the  $\chi^2$ -minimum obtained upon ument that the best fit has been obtained, it is ting procedure in some detail in order to docfurther to our confidence in the spectroscopic the curve-fitted spectra. Some of these changes interpretations rely on systematic changes in These factors have been addressed recently for Several important aspects of the spectral

amorphous materials (e.g., Walrafen, 1967; or not the minima in  $\chi'$  are regional or global McMillan et al., 1982; Mysen et al., 1982 Hartwig, 1977; Seifert et al., 1981. commonly the case for Raman spectra of fitted with symmetric Gaussian lines, as is CaAl<sub>2</sub>O<sub>4</sub>-SiO<sub>2</sub>-H<sub>2</sub>O quenched melts are best dures it was found that the spectra of hydrous nosilicate glasses. From these fitting procefitting spectra of the hydrous calcium alumiinput trial parameters were randomly varied by is not easily subjected to rigorous statistical McKeown et al., 1984; Mysen, 1990). Whether ests. However, for each fitted spectrum, the Similar procedures were employed here for 1982:

> are so small that they are statistically insignifof these aspects to the curve-fitting routine) icant (see also Mysen et al., 1982; Seifert et al ditional lines also tend to have intensities that prove the value of  $\chi^2$ . Furthermore, such adwhen additional lines do not significantly inof lines. This number of lines was taken as that ployed to determine the appropriate number served. The convergence routine is also emparameters and in the values of  $\chi^2$ . Within stavalues in order to ascertain whether the result choosing between 10 and 20 sets of differen tistical error, no variations have not been ob ing, converged fits yield different values in line 1982; Mysen, 1990, for additional discussion

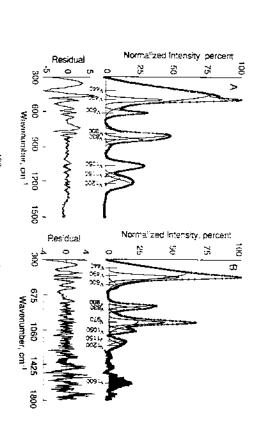
## 3. Experimental results

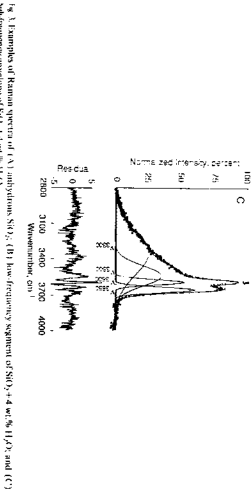
quency information summarized in Tables IItra from these two frequency ranges are shown melts on the join CaAl2O4-SiO2 are divided in Figs. 3 and 4 with detailed area and freinto two regions, Representative Raman spec-The Raman spectra of hydrous quenched

# 3.1. Low-frequency region

creasing Al/(Al+Si). The large envelope bespectra could be best interpreted in terms of al., 1982), Seifert et al. (1982) fitted four mons, 1981; McMillan et al., 1982; Seifert et trend also observed by others (Sharma and Sincreasing Al/(Al+Si) in anhydrous melts, a tween 900 and 1300 cm. becomes more inthe two bands becomes less distinct with inof which decrease slightly and the resolution of with increasing Al/(Al+Si)], the frequencies can be deconvoluted into two bands spaced segment (800-1300 cm<sup>--1</sup>) the 800-cm<sup>-1</sup> band Thorpe, 1977) those authors inferred that thee the aid of the central-force model (e.g., Sen and bands in this high-frequency envelope. With tense and shifts to lower frequency with  $\sim 30$  cm ' apart [this difference decreases (Seifert et al., 1982), in the high-frequency In spectra of anhydrous CaAl<sub>2</sub>O<sub>4</sub>-SiO<sub>2</sub> melto

Subfrequency envelope of Sitt, 1.4 wt % 11.0.





ahedra per ring and in their respective AI/ sopic studies (Sharma et al., 1981; Galeener, (Al+Si). Two three-dimensionally interconjural units that differed in the number of jetbeen suggested from other Raman spectronected ring structures in SiO, melts have also

vitreous SiO,). It was proposed (Seifert et al., multiple three-dimensional structural units in (982) that by increasing the AI/(AI+SI) in 1988, for review of the evidence suggesting (Gaskell and Mistry, 1979; see also Mysen, lution transmission electron vitreous SiO2 can also be found in high-resofen. 1983). Evidence for two structure types in microscopy

two three-dimensionally interconnected struc-1982; Phillips, 1982, 1984; Revesz and Walra-

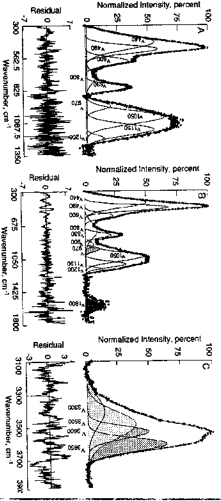


Fig. 4. Examples of Raman spectra of (A) anhydrous CA258; (B) low-frequency segment of CA258+5 wt.% H<sub>2</sub>O; (C) and high-frequency envelope of CA258+5 wt.% H<sub>2</sub>O.

alkaline-earth aluminate-silica melt, the proportion of the two structural types changed, but not the Al/(Al+Si) of the individual units (Fig. 5). This structural interpretation by anhydrous CaAl<sub>2</sub>O<sub>4</sub>-SiO<sub>2</sub> melt structure differs from that for NaAlO<sub>2</sub>-SiO<sub>2</sub> melts where Al/(Al+Si) of the individual three-dimensional structural units is positively correlated with the bulk melt Al/(Al+Si).

The low-frequency region (300–1800 cm<sup>-1</sup>) of the spectra of hydrous samples can be divided into three distinct segments (Figs. 3 and 4). In the region from 300 to ~600 cm<sup>-1</sup> all spectra show an envelope with a peak maximum near 500 cm<sup>-1</sup>. This envelope has a small shoulder on its high-frequency limb (~600 cm<sup>-1</sup>). There is a higher-frequency segment between ~800 and ~1300 cm<sup>-1</sup> consisting of a distinctive band or two bands near 800 cm<sup>-1</sup> and a group of three of five bands in the envelope between 900 and 1300 cm<sup>-1</sup>. Finally, all the hydrous samples show a distinct band near 1600 cm<sup>-1</sup>. Those of anhydrous materials do not

For all compositions, the envelope centered near 500 cm<sup>-1</sup> sharpens as H<sub>2</sub>O is dissolved in the material. Three bands have been fitted to

this envelope ( $\nu_{440}$ ,  $\nu_{490}$  and  $\nu_{600}$ ; see Figs. 3A. B and 4A, B). The central band near 490 cm<sup>-1</sup> is more dominant in the spectra of hydrous materials than in the anhydrous equivalent. Furthermore, in spectra of the hydrous samples, the  $\nu_{600}$  band is less intense relative to the rest of this frequency region (Fig. 6). This behaviour of the 600-cm<sup>-1</sup> band, first noted by Stolen and Walrafen (1976) for SiO<sub>2</sub>-H<sub>2</sub>O glass, is similar to that for melts in the system NaAlO<sub>2</sub>-SiO<sub>2</sub> (Mysen and Virgo, 1986).

appears near 970 cm<sup>-1</sup> (Fig. 3). With D<sub>2</sub>0 in crease with increasing Al/(Al+Si) (Fig. 7). In and 4). The frequencies of all the bands in the quency portion of the Raman spectra (Figs. 3 der to satisfy the spectral signal. The frequency gested that two Gaussian lines with different cm 1 (Table II). The band is slightly asymreplacement for  $H_2O$  this band shifts to  $\sim 950$ 900-1300-cm<sup>-1</sup> region show a continuous demelts results in several changes in the high-fre-H<sub>2</sub>O melts is insensitive to water content. In of the band or two bands near 970 cm<sup>-1</sup> SiO<sub>2</sub>. metric, and Mysen and Virgo (1986a) sugthe spectra of SiO<sub>2</sub>-H<sub>2</sub>O melts a narrow band line-width should be fitted to this peak in or-Solution of H<sub>2</sub>O in calcium aluminate-silica

INTERACTION BETWEEN WATER AND MELL IN CHALO, SIO5-1150

TABLE II

requencies of Raman bands in the 3(t)-1800-cm-1 range

	AI/(AI+Si)	+Si)										
	440	490	600	650	800	830	909	970	1,050	1,150	1,200	1,600
Anhy	Anhydrous:									ļ		
-	449 <u>!</u> I	495+I	608 ± F		800 1	840+I			1,062±1	1,161±1	1.2 <b>13</b> <u>1</u> 1	
0.125	453 ± 1	492±1	1 + 565		735±1	805 ± I		9521	1,069±1	1,125 ± 1	1.189±1	
	440±1	497 + 1	591 + 1		724 ! 1	803+1		954±1	1,047±1	1,145±1	$1,209 \pm 1$	
	449+	505 + 1	586 !		11111	802 + I		939 !	1.810,1	1,123 ± 1	1,197 ± 1	
0333	437±1	495 1	590 ! 1			787±1		1 i 086	1+966	1,100+1		
3 W. 6	3 WL & H2O.											
¢	439 <u>⊩</u> 1	484 F ]	589 ± 1		788 ± 1	822 ± 1		968±1	1,060±-	1,194±1	1,239 ± 1	1,621 ! !
0.125	427 + 1	490 + 1	$583 \pm 1$		728 + 3	806 ± -	914±2	967 <u>±</u> 1	1,050±1	1,167±1	1,167±1	1,627±
	409±1	499 ± 1	<b>e</b> 10 i <b>1</b>		718 ± 4	821 ± J	926 ( 5	999 ) 2	1,080+2	1,215±3	1,215±3	1,665±
0.23	456±2 461±1	508+1	594+1	630 1.5	728.1	747 ÷ 1	889 + 2	924 ± 4	1,009±1	1,207±3 1,180±1	1,207±3 1,180±1	1 : 81931
4 n/. 8	4 ml. % H2O:											
	443 1 1	480 + 1	577±2		791 L I	826+1		961±1	1.066 ± 1	$1,164\pm2$	1,221 ± 2	1,608-11
5 45. 家	S ML 186 H2O:											
£	443±1	483 ± I	570 ± 1		1 88.7	821+1		963±1	1,063±1	1,167±1	1,211+1	1,615+
تت	449 + 1	477±1	596 ± 1		712±1	803±1	902.1.2	964+2	1,052 ± 2	1,156±2	1,197±1	1,635±
1.25	440+1	\$00+E	587+		740 1 2	791 -	7.4.90K	938+	037+1	1,135 1	1,1961	1,589 1
-	454÷1	523±1	609 ± 1		725+1	800±1	890±1		0,019 ± 2	1,120±1	1,188±1	1,637±
7.5 m/	7.5 mt. % H <sub>2</sub> O:											
=	447 ± 1	484±1	578 <u>I</u> . I		788 ± 1	810±1		962±1	1,062 ±	1,16311	1,212±1	1,633 £ 1
10 mg	10 mr. % 11 <sub>2</sub> O:											
•	445 + 1	484±	572 <u>i</u> l		792±1	825±1		964±	1,064±1	1,168±2	1.215±1	1.623 !
ټڼ	449±1	490 !! 1	594+1		748 <u>†</u> 1	807 + 1	882±1	964±1	1,045 ± 1	1,131±1	1,200+1	1,629+
57	436±2 462±1	494 + I 497 + I	585 ± 1		753 + 2 696 + 2	798 ± 2 787 <u>÷</u> 1	87411	942±1	1,046+1	1,140±1 1,123±1	1.194±1 1.181±1	1,630±
\$ 84.% D <sub>2</sub> O:	0,0											
	447 1	484+]	594±1		793+1	827 ± I		948 ⊦ [	1,065 ± 1	t,174±1	1,217±1	
-	448+1	504 ± 1	593 E I		_		905 + 1	960±1	1,062±1	1,150+1	1,219 ± 1	
17.	481 + 1	216+1	500-1		7301+1	743+1	900±1	95212	1,067±5	1,00711	1,219±2	

intensity may increase in the water concentration range between 3 and 5 wt.%, and also increases slightly relative to that of the 1600-cm<sup>-1</sup> band in this water concentration range. Higher water contents (> 7.5 wt.%) may cause this intensity ratio,  $A_{970}/(A_{970}+A_{1600})$ , to de-

crease (see also further discussion on pp. 233-234).

The appearance of the 1600-cm<sup>-1</sup> band in spectra of Al-bearing melts on the CaAl<sub>2</sub>O<sub>4</sub>-SiO<sub>2</sub> join band is similar to that of pure SiO<sub>2</sub>, and also resembles that of melts on the join

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TABLEIII

Frequencies in the 3000-4000-cm | region

Laufan.	ICTCS III VIIC SIA	- 5	- Kron		Manager	Meading of Edition Indians in the S	an manana m	- 5
	Al/(Al+Si)			-		Al/(Al+Si)	•	
	3,300	3,500	3.6(X)	3,650		3,300	3,500	
3.84.6	3 ret # 1130				3.847.6	3 wt. % H <sub>2</sub> O:		i
=	3,460 ± 6	3,570 ( 3	3.595 (	3,659 + 1	=	22.8 ( 1.5	35,K+2,4	_
0.125	3,273 + 5	3,473 1 5	3,576 1 1	3,654 + 3	0.125	9.1 + 0.11	39.7 ) 4.7	
0.2	3,269 1	3,437 1 1	3,53341	3,623 + 1	0.2	2.4 (01.	$20.9 \pm 1.1$	
0.25	3.267 + 1	3,391 ( ]	3,533 ( )	3,628 + 1	0.25	23.0 / 1.1	4.5 + 0.4	
0.333	3,338 + 2	3,454 ( 2	3,387+1		0.333		38.0 / 1.8	
A WL	A wit % Hij Ch				4 1141.5	4 WL 90 H;O:		
၁	3,348 / 1	3,541 ( ]	3,595 (1)	$3.660 \pm 1$	c	45.0 / 2.4	29.30 1.4	_
3 101.1	5 wt.% 11,0:				5 144.6	5 W. M. H. D.		
0	3,305 + 1	3,515 (1	3,594 1	3,661 + 1	=	42.8 + 2.5	30.9 + 1.4	_
0.125	3,273 + 1	3,487 ! 2	3,565+1	3,641 + 2	0.125	5.4 / 0.8	47.4 1 2.5	٠
0.2	3,412 / 9	3,498 ( 3	3,546 1	3,623 + 1	: =	13.9 (1.5	35.8   7.3	
0.333	3,305+2	3,505 ( )	3,568 ( 2	3,634 + 2	0.133	27.3 + 1.6	48.2 ) 1.6	_
7.5 H	7.5 80.% H <sub>2</sub> O:				Z.5 w	7.5 m; 4 H; O:		
=	3,307 + 1	3,514 ( )	3,595.1	3,658 ). (	0	39.3 ( 1.3	44.8 ) 1.3	
10 m	10 wa. % 115O:				10 H	10 wt. % H2O:		
5	3,317+2	3,505 ) )	1.595 ) [	3,657+1	=	55.4 1 1.7	30.8 / 1.0	
0.125	1.282 1.6	3,480 - 6	3,570+2	3,660 1	0.125	14.6 10.7	21.6   0.8	
0.25	3,271 1 9	3,434 + 4	3,586 11	3,629 1 1	0.25	24.0 + 2.2	29.3 + 1.4	_
5 W. S	S 42. 46 DzO:				5 HZ.1	5 ur. % D <sub>5</sub> O:		
\$	2,536 + 1	2,578 + 4	2,642 + 2	2,700 + 1	=	42.4 + 3.5	10.74 L.8	
0.125		2,552 ! 2	2,683 ( 2	2,683 1 1	0.125		59.8 ! 4.8	2
0,2	2,205 + 1	2,501 (1	2,62	2,692 + 1	0.2	%.I + 1,2	37,4 ± 2.1	
0.25	2,215 ( 3	2,480 + 3	2,601 ) [	2,679 + 1	0.25	4,410,9	44.1 ) 2.8	4
								1

presence of molecular H<sub>2</sub>O in the sample signed to H-O-H bending and requires the Mysen and Virgo, 1986a). The b (Scholze, 1960; Stolper, 1982). NaAlO<sub>2</sub>-SiO<sub>2</sub> (McMillan and Rem

mass difference of the oscillators (Freund, anticipated frequency decrease caused by the substitution of <sup>2</sup>H for <sup>3</sup>H is consistent with the OH stretching (e.g., Stolen and Walrafen, 1976). The ~15-cm<sup>-1</sup> frequency reduction by quenched hydrous silica melt is assigned to Si-The 970-cm band in the spectra of

3,659   1   3,659   1   1   3,654   3   1   3,628   1   1   1   3,641   2   1   1,641   2   1   1,643   1   1   1   1,643   1   1   1   1   1   1,643   1   1   1   1   1   1   1   1   1	# Relative  # wt.#6  0.1125 0.2 0.233  # wt.#6 0  5 wt.#6 0 0.125 0.125	### ##################################		3,600 3,600 12,3+0,8 36,7+4,3 36,7+4,9 59,4+1,9 59,4+1,7 11,7+0,6 15,9+0,6 21,4+3,5	3,650 3,650 29,3117 12,7113 47,5109 13,310,6 13,310,8 13,310,8	### Weard individual by ###################################	mil bands in the miles of the first of the f	6000 6000 50 (0) 52 (0) 52 (0) 52 (0) 53 (0) 50 (0) 57 (0) 57 (0) 57 (0)	Min (190) 6 Min (190) 6 Min (190) 7 Min (	800 cm   range 600 600 600 600 600 600 600 600 600 60	(11 - 11 - 12 - 12 - 12 - 12 - 12 - 12 -
3,660+1			29,3+1,4	11.7 + 0.6				41,9 + 4.1 19.6 + 1.3 8.2 ( 1.3 10.9 + 2.1 79.8 + 1.6	41,9 + 4.1 19.6 + 1.3 8.2 ( 1.3 10.9 + 2.1 79.8 + 1.6	79.01/02 10.0 4.1 [21   1.3   10   0.3   10.6 (1.1   18.5   1.7   5.0   0.3   10.6 (2.1   18.5   1.7   5.0   0.3   10.6 (2.2   10.8   1.7   5.0   0.3   10.9 (2.2   2.3   4.4   4.5   3.1   0.3   20.8 (1.0   6.2   0.7   5.7   0.3	7941702 41.9 44 121 2.3 10 4 0.3 196 4 (1) 18.5 2.7 50 2.6 8.2 0.2 408 13.7 54 4 4 4 5 10.9 2.2 23.4 4 4.5 3.1 0.8 10 11 29.8 11.0 6.2 20.7 57 20.4
3,661 F1 3,641 F2 3,623 F1 3,648 F1 3,634 F2	0 0.125 0.2 0.25 0.133	42.8 ± 2.5 5.4.± 0.8 13.9 ± 1.5 19.1 ± 1.5 27.3 ± 1.6	30.9   1.4 47.4   2.5 35.8   7.3 35.9   2.2 48.2   1.6	15.9 ± 0.6 37.5 ± 2.0 21.4 ± 3.5 42.9 ± 2.2 12.7 ± 0.8	13,9+0.6 13,9+0.6 28,9+5.9 2.1.1-0.2 11.8+0.8		4 m m H/O; 32 (0.3	%H,O: 32 (113) %H,O:	%H,A; 3,2 (1),3 = 9,0 (4),7 %H,A;	%H,O; 3.2 (1)3 9,0 (4);? %H,O;	% H/A 32 (1)3 - 9,0 (4)3 - 2 ( (4)) % H/A
3,658 ). [	7.5 x 0	7.5 wt.% H <sub>2</sub> O; 39.3 + 1.3 10 wt.% H <sub>2</sub> O;	44.8 ) 1.3	%.T = 0.4	7.9 1 0.4		20.63.21 22.39.14 22.39.14 23.18.44.18.6		22.0 F 2.1 22.0 F 1.4 14.4 F 10.8	22.0 F (4 13.2 ) 6.2 22.0 F (4 13.2 ) 6.2 14.4 + 0.6 15.0 + 0.6 13.0 + 0.8 19.7 + 1.7	22.0 F (4 13.2 ) 6.2 22.0 F (4 13.2 ) 6.2 14.4 + 0.6 15.0 + 0.6 13.0 + 0.8 19.7 + 1.7
1 3,657+1 12 3,660+1 14 3,633+1 14 3,629+1	0 0.125 0.2 0.25	55.4   1.7 11.6   0.7 19.7   1.1 24.0   2.2	30.8 ) 1.0 48.3 + 2.5 21.6 + 0.8 29.3 + 1.4	7.6 ± 0.3 30.4 ± 2.0 54.8 ± 1.7 44.7 ± 4.5	6.2+01 9.7+0# 3.9+01 2.0+61		"Swith H/O. 364 + 2,0		11.2 + 13.7	11.2 + 13.7	11.2007 25104
2,700   1 2	5 Hz. 0 0.125 0.2 0.25	5 HZ % D <sub>2</sub> O; 42.4+3.5 125 8.1+1.2 25 4.4+0.9	10.74 L.8 59.8 i.4.8 37.4±2.1 44.1 J.2.8	25.1 ) 2.4 28.4 ) 3.0 45.5 ) 2.3 42.5   2.7	21.8+1.7 12.0+1.4 9,0+04 9,0+04		31,3 11,7 115 31,4 11,8 2 20,9 11,1 25 24,2 11,2 1 96,5 10,6	31,3 11,7 990 06 125 314 11,8 32 165 2 23,9 11,3 16,9 166 25 24,2 11,2 8,2 113,5 1 60,9 13,42	113 113 314 118 209 113 242 112	H.3+1.7 9,9+0,6 H.4+1.8 8,2+0,5 23,9+1,3 8,3+0,6 24,2+1,2 8,2+0,8 4,2+1,2 8,2+0,8	H.3+1.7 9,9+0,6 H.4+1.8 8,2+0,5 23,9+1,3 8,3+0,6 24,2+1,2 8,2+0,8 4,2+1,2 8,2+0,8
nmele, 1986; band is as- requires the	1982 cal 17	) in SiOI 2-cm de	[982] in Si-OH and Si-OD (with a theoretical [2-cm] decrease, see also additional discussion on p. 232). Mysen and Virgo (1986a)	D (with a also addit	theoreti- ional dis-		;	366 (2.0 8.8) (4.6) (125 22.5) (1.2) (2.6) (4.9) (4.5) (1.5) (2.5) (4.5) (1.5) (2.5) (4.5) (1.5) (4.5) (1.5) (4.5) (1.5) (4.5) (1.5) (4.5)		8.81-0.6   17-0.3 	8.8 1 0.6 15.6 1 0.9 14.5 1 1.15 9.0 1 0.5

ert et al., 1988). Whether more than one Siof SiO<sub>2</sub>-H<sub>2</sub>O glasses (Farnan et al., 1987; Eckone Si-OH bond in Si-O(OH) tetrahedra in suggested that the two bands fitted to the 970tetruhedra in SiO<sub>2</sub> (Yamamoto et al., 1983) analogy with multiply F for O-substituted SiO, cm - region of the SiO2-H2O spectra could be cussion on p. 232). Mysen and Virgo (1986a) This suggestion is not supported by 'H NMR interpreted as being due to perhaps more than

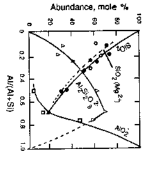
A HIN

Al/(Al+Si)	() X							:			
i.	\$	Ē	<u> </u>	CICIK	*	∰ Egy	1978	0.00	1,150	1,2100	1,618
Supply by the state of the stat									:		:
44.44.74	-										
13 EL 21		50104			2.4104	1.001.001		0 8 1 10 /	6,3 4 0,7	470-117	
		7 7 0 K		77.57		_		70.4 7 7.1			
		10 1 Kill		£ .	50.00	_	41101	200		2 N 1 0.2	
_		10405		3,1101	1		20.02	E1 1871	249113	3.77 0.3	
$O^{H(\mathbb{R}, \mathbb{M})}$											
	12101.3	10101		90+15	7.1 II.		153+13	44-	111	-	
-		20.507		22.404	¥1.0)	top	.710.4	20 6 1 1 2	34 189	1.1	4.0
		F4 1417		4.91.14	6.1 × 0.8		F. 1 - 1 - 2	5 1		45113	
07 1 8762 331 17 1 670 1 1		F3 - 4.5 KD + 1.6	-	X =	22 F0.7		5.3 1 2.4	27) 17.4	11.9 + 2.6	2,010,8	37104
Aut & HyO:								;		:	
32 (10)	9,014,02	?   · #:		444113	31 HH		<del>s</del>	S 184	5.5 1 @ 5	22103	5.0 1.03
Of H & the											
	12.5 1 (1.7	1919		\$ 105	4.8 10.4		22.0116	<u> </u>	10.12	7	÷
7	W ( )	11114		2.0 1 # 5	1 th 1 K'z	÷ - + -	1.5 1 0.4	<b>1.0</b>	1.0 1 0.9	27101	
2 144414		104		71104	1.0.1	1.0 1.0.1	11.57	77.1561	9.4 ( 0.7	2.5   0.2	58105
_	19711.	40103		941 36	0.00 L RG	# - <del>*</del> <del>*</del> - <del>*</del> <del>*</del> - <del>*</del> - <del>*</del> - <del>*</del> - <del>*</del> - <del>*</del> - <del>*</del> <del>*</del> - <del>*</del> - <del>*</del> - <del>*</del> - <del>*</del> - <del>*</del> - <del>*</del> <del>*</del>	5.8 1 0.4	29.7 F 1,0 28.9 F 1,7	29 - 103 170 - 164	2.9 (0.1	2 % 2 %
O'H W. Tan S.											
36,4 + 2,0	36.4 + 2.0 - 11.2 + 0.7	53104		19145	80 - P.00		#.j + j.w	7410.5	3.7 ( )3.4	4,0 + 0,6	12.5 4 D.4
10 H 1 1 11 20:											
11.3   1.7	9,010,00	1,000		67101	<u>-</u>		·	<b>*</b>			-
WILLIAM SE	K31163	11 11		1111	7 7 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		7 - 1.0				
23.9 F.1.3	83406	5.11114		# **	<u> </u>			11.0	0.14.07	10.7	4.4
24.2 + 1.2	8.2 1 12.5	(J + D.)		9.0 ( 0.5	3.8   6.3	1.19   87		30.7 1.5	6.1 10.4	26192	93 (85
$m \in D_i O_i$											
	970 L R'R	1700.1		4.3 t U v	4.8 10.1		111-3 1-41 7	\$6.04	· ·		
	15.6 ( 0.9	10144		X 2 + D 4		17:01				104	
23.4 1 1.8	14.5 1 1.15	D.N. I O.S.		6.0 1 (0.74 1 0.4		E 20			25   05		
	100				K.0 + 0.4		. D.1	2.0 ' 0.2		%.5 × 1.#	1,4 + 0.2
								ļ			:

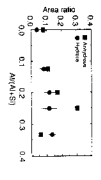
melts (or glasses) remains open to further investigation. Oll bond may exist in SiO. H.O quenched

sty in this general frequency region. This in-Fig. 3). However, there is an increase in inten-970 cm 1 in contrast to SiO, 41,O mells (see ples there is no visually well-defined band near In the spectra of CaAt,O,-SiO,-H,O sam-

D<sub>i</sub>O (Figs. 8 and 9; see also Table II). capily in frequency as H<sub>2</sub>O is substituted with Page band (or bands) does not shift signiff-970-cm. band in hydrous silica glasses, the (Al+Si). In contrast to the behavior of the 900 cm  $^{-1}$  ( $\nu_{900}$ ), depending on the Al/ 960-cm ' (also denoted  $p_{070}$  in this text) and tensity results in one or two bands fitted near



in anhydrous CaAl2O4-SiO2 melts as a function of Al/ Fig. 5. Abundance of three-dimensional structural units (Al+Si) (data from Seifert et al., 1982)



function of their AI/(AI + Si). v<sub>600</sub> + v<sub>440</sub> + v<sub>490</sub> for hydrous and anhydrous melts as a 6. Area of  $\nu_{600}$  band relative to the area of

curs in hydrous melts on the join NaAlO2-SiO2 cm<sup>-1</sup> ( $\nu_{900}$ ) as shown, for example, in Fig. 4B one near 970 cm<sup>-1</sup> ( $\nu_{970}$ ) and one near 900 Virgo, 1986a) and has also been reported in (McMillan and Remmele, 1986; Mysen and 8A). A similar Raman band near 900 cm<sup>-1</sup> ocof deuterium for hydrogen in the water (Fig. the frequency of the  $\nu_{900}$  band and substitution ples, the band can be deconvoluted into two, (Silver and Stolper, 1989). infrared spectra of quenched melts on this join There are no systematic relationships between In the spectra of the most water-rich sam-

gen (Mysen and Virgo, 1986a). However, OH stretching, In the latter reports, its lower 900-cm<sup>-1</sup> band could be assigned to (Si,Al)-Silver and Stolper (1989), suggested that the cm-1 band was assigned to nonbridging oxyfrequency compared with the 970-cm<sup>-1</sup> Si-OH McMillan and Remmele (1986), as adopted by In the NaAlO<sub>2</sub>-SiO<sub>2</sub>-H<sub>2</sub>O melts, the 900-

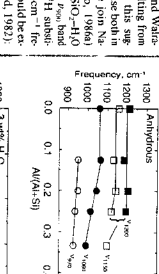
> pected from the relationship (Freund, 1982): quency reduction of the  $\nu_{von}$  band would be exstretch band in SiO2-H2O (Stolen and Walra tuted for 'H in the water. An  $\sim 10$ -cm<sup>-1</sup> fredoes not vary as predicted when 2H substihere (Table II), the frequency of the  $u_{900}$  band and those observed for CaAl<sub>2</sub>O<sub>4</sub>-SiO<sub>2</sub>-H<sub>2</sub>O AlO<sub>2</sub>-SiO<sub>2</sub>-H<sub>2</sub>O (Mysen and Virgo, 1986a) the Raman spectra of melts on the join Nagested assignment is not likely because both in the Al=Si substitution. However, this sugfen, 1976) was rationalized as resulting from

 $\nu^{\text{SiOH}}/\nu^{\text{SiOD}} = (m^{\text{SiOD}}/m^{\text{SiOH}})^{1/2}$  $\widehat{\Xi}$ 

to the conclusion that the 900-cm -1 band is not cal to that by Mysen and Virgo (1986a), leads same concentration of D2O in substitution for OH bond, the ratio is 1.0113. If the P900 band and m = oscillator mass. For an equivalent  $\Lambda$ 1ent melt compositions. bridging oxygen is the assignment in the pres interpretation that the band is due to nondue to (Si,Al)-OH, Therefore, the alternative frequency shift. This analysis, which is identi-H<sub>2</sub>O. However, there is no evidence of such a the same band should be at 890 cm<sup>-1</sup> with the is due to (Si,Al)-OH stretching at 900 cm where  $\nu = \text{frequency}$ , with  $\nu^{\text{Sion}}/\nu^{\text{Sion}} = 1.011$ ;

ception of the CA2S4 composition melt [Al/ can be explained as the result of increased Al/ of these bands with either  $H_2O$  or AI/(AI+Si)relationships to H<sub>2</sub>O concentration and Al/ AlO2-SiO2 melts, exhibits similar functional which was not observed in the hydrous Nawith increasing water content. The  $\nu_{900}$  band, observed (Table II). This behavior differs from (Fig. 8) at constant Al/(Al+Si) with the exerally decrease with increasing water content to the  $\nu_{970}$  and  $\nu_{900}$  bends. the 900-cm<sup>-1</sup> band in NaAlO<sub>2</sub>-SiO<sub>2</sub> where (Al+Si)=0.333] where the opposite trend is (Al+Si) in the structural units that give rise (Al+Si) (Fig. 9). The frequency decrement there is a systematic increase in its frequency The frequencies of the 900-cm<sup>-1</sup> band gen

1150- and 1200-cm<sup>-1</sup> bands together with their The intensity relations among the 970-



Frequency, cm<sup>-1</sup>

9

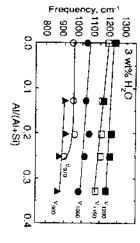
ร์ ร์

1150

Al/(Al+Si)

1300

10 wt% H<sub>2</sub>O

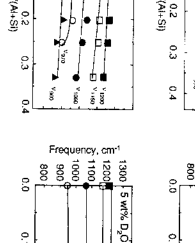


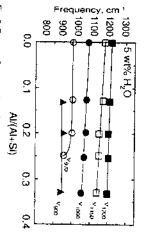
0.1

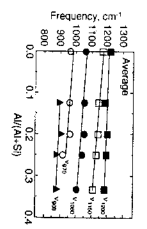
0.2 Al/(Al+Si)

0.3

1080



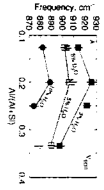


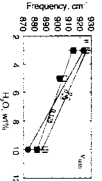


a function of AI/(AI+Si) at total water concentrations indicated. The average values are for all the  $H_2O$  contents (the Fig. 7. Raman frequencies of important bands in the high-frequency segments of the low-frequency portion of spectra as samples with D<sub>2</sub>O are not included)

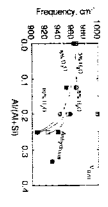
sionally interconnected network. The  $\nu_{970}$  band part of the signature of the three-dimendrous calcium aluminate-silica melts (Mepossible assignments of the 970-cm | band; est because there are at least three different Millan et al., 1982; Seifort et al., 1982) that is (Al+Si) and H<sub>2</sub>O concentration are of interfrequency relationships as a function of Al/ (1) There is a band near 970 cm. in anhy-

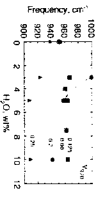
vs. hydrous CaAl<sub>2</sub>O<sub>4</sub>-SiO<sub>2</sub> melts it would apferent behavior of the \$\nu\_{970}\$ band in anhydrous pear that the  $u_{970}$  band in hydrous calcium al-Al/(Al+Si) (Fig. 10B) in particular with the lowest water contents (3 wt.%). From this difin hydrous melts, the area ratio, insensitive to  $\Lambda I/(AI+Si)$  (Fig. 10A), whereas  $(A_{970} + A_{1150} + A_{1200})$ , tends to increase with intensity in anhydrous CaAl<sub>2</sub>O<sub>4</sub>-SiO<sub>2</sub> melts is A970/





(Al + Si) at constant  $H_sO$  content (A) and as a function of total water content at Al/(Al + Si) (B) indicated on lines. Shaded squares are from D<sub>2</sub>O samples. Fig. 8. Frequencies of  $\nu_{\text{vor}}$  band as a function of Al/

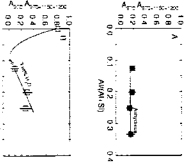


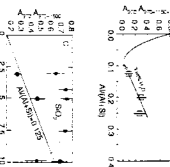


to 3 wt.%  $H_3O$  are dashed as it is not likely that the  $\nu_{sm}$  band in spectra of anhydrous samples is the same band as of total water content at AI/(AI+Si) indicated on lines. Fig. 9. Frequencies of  $\nu_{\rm trit}$  band as a function of Al/ (Al+Si) at constant H<sub>2</sub>O content (A) and as a function Shaded squares are from D<sub>2</sub>O samples. The lines from 0

near 970 cm. I in the anhydrous equivalents. uminosilicate melts is not the same band as that

(2) As observed in the spectra of quenched





*shaded* for comparison. Al-bearing SiO<sub>2</sub>-H<sub>2</sub>O quenched melt in (C) is shown us as in the Al-bearing samples. The  $A_{ijn}/(A_{ijn} + A_{inin})$  for dashed arrow as it is not likely that this band is the same data point for AI/(AI+Si) in (B) is connected with Fig. 10. Selected area ratios involving the  $\nu_{\nu_{Al}}$  bands as a function of H<sub>2</sub>O contents for compositions indicated. The

H,O. ₩1%

samples (Fig. 10C). Finally, in SiO,-H<sub>2</sub>O increase with water content in the aluminous with the highest water content, this area ratio melts the  $A_{070}/(A_{070}+A_{1600})$  is insensitive to ble V). Furthermore, whereas in SiO<sub>2</sub>-H<sub>2</sub>O higher in the Al-free (SiO2-H2O) melts (Ta- $(A_{970} + A_{1150} + A_{1200})$ (where it is assigned to Si-OH stretching) dif-Stolen and Walrafen, 1976). The behavior of stretch vibrations also are near 970 cm <sup>-1</sup> (e.g., melts in the system SiO<sub>2</sub>-H<sub>2</sub>O (Fig. 3), Si-Ol H<sub>2</sub>O concentration and may actually decrease  $(A_{vm} + A_{tolio})$  area ratios are significantly in two important ways. Both the  $A_{\rm out}$ fers from its behavior in Al-bearing materials the 970-cm<sup>--1</sup> band in Al-free SiO<sub>2</sub>--H<sub>2</sub>O melts and <del>f</del>e Augus

> stretching. the atomic wright of hydrogen was observed, reconcile with an assignment to (Si,AI) .OH hydrous  $CaAl_{2}O_{3}$ -SiO<sub>2</sub> melts is difficult to Al-bearing melts, no frequency dependence of sensitive to 'H substitution for 'H, whereas in melts the frequency of the 970-cm 1 band is Therefore, the behavior of the Para band in the

oxygen) formed in these originally fully polymerized melts through interaction with H<sub>2</sub>O. lects (Si,AI)=O (O denotes nonbridging

the the 300-1800-cm | spectral region alone certained from the high-frequency segments of sents the correct interpretation cannot be ashand. Whether alternative (1) or (2) reprepolymerized structured unit that contributes plained by increasing AI/(Al+Si) in the decreasing frequency with increasing Al/ (aAl<sub>2</sub>O<sub>4</sub>-SiO<sub>2</sub> metts. For example, the deband is controlled by structural features that (in part or in full) to the behaviour of this (Al+Si) in the hydrous melts could be exare absent in SiO.-H.O and in anhydrous in hydrous calcium aluminosilicate melts the and H<sub>2</sub>O concentration; or (2) alternatively as a function of Al/(Al+Si) and H.O content Off or (Si. Al)-OH bonding the contribution nonbridging oxygen. The intensity variations  $(SiAI) = O^n$ , or both, varies with AI/(AI + Si)from (Si,Al)-O relative to either Si OH or lead to the suggestion that; (1) if there is Siat least to a degree has a contribution from SiO,-H<sub>2</sub>O on the other, it appears reasonable (aAl,O<sub>4</sub>-SiO, melts on the one side and with that the  $\nu_{vm}$  band in CaAl<sub>2</sub>O<sub>4</sub>. SiO<sub>4</sub>-H<sub>2</sub>O melts In light of the differences with anhydrous

# 32. High-frequency region

Il stretch vibrations, the frequencies of which are sensitive to the electronic properties of the bands in this envelope are the fundamental Oem 1) is broad and asymmetric with a tail toward lower frequencies (Figs. 3C and 4C). The The high-frequency envelope (3000-4000

(3) Finally, it is possible that this band re-

tached and to hydrogen bonding, metal cation to which the OH group is at

a two-atom group (Langer and Latturd, 1980) ship between oscillator mass and frequency to OD stretching bands result from the relation samples occurs between 2000 and 3000 cm The frequency differences between the OH and The equivalent envelope for deuterated ;

$$\nu = (f/2\pi c)^{1/2} (1/m_1 + 1/m_2)$$

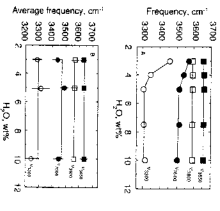
 $v_{\rm ero} = 1.37$ . From this relationship, a maxicorresponds to 2600 cm. 1 for OD, mum Raman intensity near 3600 cm. 1 for OH 1980), that the frequency ratio,  $p_{\rm OH}$ found (Hartwig, 1977; Langer and Lattard r is the speed of light; and  $m_1$  and  $m_2$  are the where  $\nu$  is the frequency; f is the force constant better resolved than those from the Al-bearing atomic masses of the oscillators. It has been The spectra from the SiO2-H2O melts are

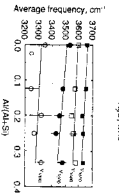
only one sharp band with a shoulder on its highcerned visually near 3600 cm 1 (Fig. 3C). In frequency side. alummous samples, visual inspection reveals samples and typically two peaks can be dis-

and 116 + 26 cm 1 for SiO, and 128 + 33 and and  $\nu_{3800}$ ) are considerably greater (197 ± 66 over all H<sub>2</sub>O and Al contents). The FWHHsystem SiO<sub>2</sub>-H<sub>2</sub>O<sub>4</sub> the frequencies of the P<sub>4,000</sub> ues of the two lowest-frequency bands (Frum values of the two highest frequency bands in than those of the  $\nu_{total}$  and  $\nu_{total}$  bands. In the 109 + 36 for Al-bearing samples, respectively) of the H<sub>2</sub>O-bearing materials (80 ± 13 and the deuterated samples are comparable to those Phon and Physic bands, respectively, averaged are somewhat broader in Al-bearing samples water-bearing silica glasses). These two bands cm ', respectively (averaged over all the sharp with full widths at half height (FWHH) 39 / 2 cm 1, respectively), The FWIIII-val-(FWHH is  $82\pm21$  and  $48\pm14$  cm  $^{-1}$  for the for the Al-free samples of 33±6 and 34±3 frequency bands ( $\mu_{\text{total}}$  and  $\mu_{\text{total}}$ ) are quite bands litted to all the spectra. The two highest The best fit is generally obtained with four

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of  $\nu_{3600}$  and  $\nu_{3650}$  (Fig. 11C), as also found for slight, but distinctive decrease in v 1600 and v 3650 correlated with Al/(Al+Si) (Fig. 11C) with a ever, all the band frequencies are negatively what more sensitive to Al/(Al+Si) than those nous. The  $\nu_{1300}$  and  $\nu_{1500}$  frequencies are some hydrous, aiuminous samples (Fig. 11B). Howhydrous melts in the system  $NaAIO_2-SiO_2$ frequency as the samples become more alumi tional relationship between the \$\nu\_{3300}\$ and \$\nu\_{3500}\$ (Mysen and Virgo, 1986a). frequencies and water content in the spectra of (Fig. 11A). In contrast, there is no clear funcand  $\nu_{3500}$  bands decrease with water content





shown when error is less than the symbol size standard error of the average values. Error bars are not as a function of Al/(Al+Si) (C). Error burs represent content (B), and average values of all Al-bearing samples values for all Al-bearing samples as a function of H<sub>2</sub>O  $SiO_2-H_2O$  as a function of water content (A), average Fig. 11. Frequencies of fundamental OH stretch bands in

crease in intensity of the two highest-frewater content (Figs, 12 and 13). quency bands  $(\nu_{1600}$  and  $\nu_{1680})$  with increasing increase of the two lowest-frequency bands observation results from the general intensity (Figs. 3C and 4C). For all compositions, this rapid growth over a broad frequency region < 3600 cm<sup>-1</sup> as the water content increases Visual inspection of the the spectra reveals a and  $\sigma_{1500}$ ) and a complementary de-

velope in  $H_2O$  samples ( $\nu_{33000}$ ,  $\nu_{35000}$ ,  $\nu_{36000}$  and to the four bands in the 3000-4000-cm<sup>-1</sup> en spectral characteristics and may be equivalent  $\nu_{3650}$ , respectively]. Thus, the four bands in the which data have been obtained (0-0.25) for age  $v^{OH}/v^{OD}$  in the Al/(Al+Si) range  $\nu_{2300}/\nu_{3300}, \nu_{2500}/\nu_{3500}, \nu_{2600}/\nu_{3600}$  and  $\nu_{2690}$ the bands are 1.39, 1.36, 1.36 and 1.35, for tra relate to the four bands from the H<sub>2</sub>O-bear- $(\nu_{2000}, \nu_{2500}, \nu_{2600})$  and  $\nu_{2650}$  have the same 2000-3000-cm<sup>-1</sup> envelope for D<sub>7</sub>O samples to the theoretically predicted value [the avering samples with frequency ratios quite similar in the high-frequency portion of the  $D_2O$  spec From eq. 2, the frequencies of the four bands

bonding will lower the O..H bond strength of the two lowest-frequency bands in SiO2and (2) the relative intensity of the  $\nu_{3300}$  and Systematic frequency reduction of OH stretch the total water content is increased. Hydrogen H<sub>2</sub>O is consistent with greater importance of creasing H<sub>2</sub>O concentration (Figs. 11A and v<sub>3500</sub> bands increases significantly with inthat: (1) the frequencies of the  $\nu_{3650}$  and  $\nu_{3600}$ OH groups from Si-OH bonds (P3650 and hydrogen bonding among water molecules as The frequency decrease with H<sub>2</sub>O content bands are independent of H<sub>2</sub>O concentration assignment is consistent with the observations ica melts can be assigned to OH stretching in  $\nu_{3600}$ ) and from OH groups in molecular H<sub>2</sub>0  $(\nu_{3300}$  and  $\nu_{3500}$ ). In the SiO<sub>2</sub>-H<sub>2</sub>O system, this The four bands in the spectra of hydrous sil-

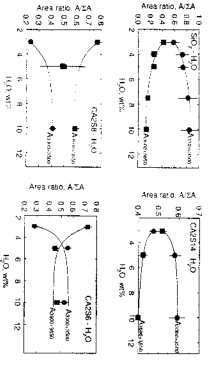


Fig. 12. Relative area of the two low-frequency ( $\nu_{\text{turn}}$  and  $\nu_{\text{tsto}}$ ) and the two high-frequency ( $\nu_{\text{turn}}$  and  $\nu_{\text{tsto}}$ ) bands daine to the complete scattering intensity of the OH stretch band envelope as a function of H<sub>2</sub>O content for composimass indicated,  $\sum A = A_{13000} + A_{33000} + A_{36001} + A_{3600}$ 

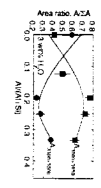
drogen bonding SiO2-H2O metts with water concentration bond strength lowering the OH stretch force (Fig. 11A) may be due to the influence of hy- $\nu_{\text{two}}$  and  $\nu_{\text{3500}}$  bands in the spectra of quenched comparatively rapid frequency reduction of the constant and, therefore, the frequency. The increased hydrogen bonding lowers the OH droxyl-bearing minerals (Dobson et al., 1989). 1955), as also recently demonstrated for hybonding is well documented (Nakamoto et al.,

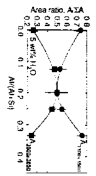
(Al+Si) of the melt (Fig. 11B) dent of total H<sub>2</sub>O content regardless of AI/ (3) in contrast to the SiO<sub>2</sub>-H<sub>2</sub>O system (Fig. sgnificantly broader than in SiO,-H<sub>2</sub>O; and (Fig. 11C); (2) the  $\nu_{3650}$  and  $\nu_{3600}$  bands are crease with increasing Al/(Al+Si) of the mel materials: (1) the frequencies of the  $\nu_{M50}$  and differences. In the spectra of the Al-bearing those in SiO<sub>2</sub>-H<sub>2</sub>O, there are some important IIA), all the band frequencies are indepenrum bands show a small, but systematic desty trends in aluminous samples resemble Although the overall frequency and inten-

The Passe Passe and Passe band frequencies and The negative correlation (Fig. 11) between

One cannot rule out some (Si,Al)-OH in these  $v_{970}$  band is reduced by  $\lesssim 5$  cm<sup>-1</sup> (Fig. 9). nous samples. The observed frequency of the signed to (Si,Al)-OH stretching in the alumicm "1; see Table II) if the P970 band was as fitted line positions (typically  $\pm 1-4$  cm  $^{-1}$ ). which is barely outside the uncertainty in the in SiO<sub>2</sub>-H<sub>2</sub>O vs. SiO<sub>2</sub>-D<sub>2</sub>O  $A\nu^{D/H} = 15 \pm 3$ did not, however, show the expected frestretching. For example, the 970-cm | band also has a contribution from (Si,Al)-OH reduction is expected from eq. 1 (as observed for hydrogen (Fig. 9),  $A \sim 13$ -cm<sup>--</sup> frequency quency-dependence of deuterium substitution sion on pp. 233-235) unless this latter band cult to reconcile with the behavior of the Popul band in CaAl<sub>2</sub>O<sub>4</sub>-H<sub>2</sub>O melts (see also discus-However, this interpretation somewhat diffi-(Si,Al)-OH or Al-Oh bonds in the samples from substitution of Al3+ for Si4+ to form fleet a weakening of the O-H bonds resulting bands with increasing Al/(Al+Si) might re-The frequency reduction of the  $\nu_{3650}$  and  $\nu_{3600}$ ion (M) in possible M-OH bonds in the melts ing the electronic properties of the metal cat-Al/(Al+Si) may be rationalized by consider-

3





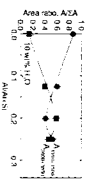


Fig. 13. Relative area of the two low-frequency  $(\nu_{\text{tim}}, \text{and} \nu_{\text{tim}})$  and the two high-frequency  $(\nu_{\text{tim}}, \text{ind} \nu_{\text{too}})$  brinks relative to the complete scattering intensity of the OH stretch band envelope as a function of  $\Delta t/(\Delta t + Si)$  for the H<sub>2</sub>O concentrations indicated.  $\sigma \Delta = A_{\text{tim}} + A_{\text{too}} + A_{\text{to$ 

melts, but the contribution from such bonds must be comparatively small. With regards to the spectral topology in the 3000-4000 cm<sup>-1</sup> region, it is suggested that a small fraction of the structurally bound OH could be a mixed (Si,Al)-O(OH) tetrahedral environment, or there may be mixtures of Si-OH and Al-OH, where the frequencies of the respective OH stretch bands in the 3600-cm<sup>-1</sup> envelope could not be resolved with the fitting procedure. The considerably greater line width of the  $\nu_{\text{toto}}$  and  $\nu_{\text{toto}}$  bands in spectra of the aluminous could be interpreted as consistent with the latter suggestion.

The two bands lower-frequency bands ( $\nu_{\text{TMO}}$  and  $\nu_{\text{MMO}}$ ) could be due to OH stretching in molecular water. Their increased intensity with increasing water content is consistent with an

et al., 1988). The presence of molecular H<sub>2</sub>0 are isolated. the H<sub>2</sub>O molecules in aluminositicate glasses gestion is consistent with 'H NMR informaof water present (Paterson, 1982). This sugzation in the melt does not vary with amount tent (Table II) suggests that water polymen-1600-cm 1 band is insensitive to water con III). The observation that the frequency of the CaAl,O<sub>4</sub>-SiO<sub>2</sub> (Fig. 4B; see also Tables II and bending mode, with a frequency near 1600 is also consistent with the distinctive H-O-H NMR spectroscopy; e.g., Stolper, 1982; Ecken samples (as inferred from infrared (IR) and increased abundance of molecular H<sub>2</sub>O in thes tion (Eckert et al., 1988) which indicates that t, in all hydrous samples from the join

contributions from OH groups bonded to other compositions. An alternative is that in addslightly with Al/(Al+Si) (Fig. 11C). The freof the  $\nu_{1300}$  and  $\nu_{1300}$  band is > 50% of the enequivalent to 1.35 wt.% H<sub>2</sub>O [a water concenjob Na(OH)-SiO, with NaOH concentration metal cations (Ca21, or Al11, or both). This groups in H<sub>2</sub>O, this frequency region also has tion to some intensity contribution from OH frequencies, which is not observed for these water concentration dependence of the band SiO<sub>2</sub>-H<sub>2</sub>O (Fig. 11A) one would also expects become more aluminous. If so, in analogy with quency shift could be interpreted to result from (Fig. 11A and B), In the latter compositions crease with increasing water concentration quenched melts where their frequencies de SiO, melts differs in that in SiO,-II,0 bands in hydrous silica and hydrous CaA,O,molecular H<sub>2</sub>O present (Mysen and Virgo spectroscopy indicate a significant amount of tration range whether neither IR nor Raman increased hydrogen bond strength as the melo the  $\nu_{\rm AMD}$  and  $\nu_{\rm AMD}$  band frequencies decrease whereas those from CaAl<sub>2</sub>O<sub>4</sub>-SiO<sub>2</sub>--H<sub>2</sub>O do not 1986b; Silver and Stolper, 1989), the intensity (Mysen and Virgo, 1986b) that in melts on the interpretation is consistent with observations However, the behavior of the Pring and Prin

> not be resolved with the fitting procedure. spectrum, but that the individual bands could the melts and contribute to this portion of the schosen, no more than 0.5% of the species is ofOH (H...OH, AL..OH and Ca...OH) exist in ton to H-OH. It is likely that all three forms from both AI=OH and Ca=OH bonds in addimay have contributions from OH stretching partion of the two low-frequency intensities selope. Thus, there is evidence that at least a molecular  $H_2O_s$  the  $p_{som}$  band also represents regardless of which model of water speciation centration equivalent to 2.5 wt.% H<sub>2</sub>O where, near 50% of the intensity in the OH stretch enbre OH stretch envelope. In the system SiO,-Al(OH), with Al(OH), with Al(OH), com-

#### 4. Discussion

HO) may depend on the conditions of tembe results (Dingwell and Webb, 1989a, b). or example, could affect the interpretation of resture quenching where the quenching rates. shown in Fig. 2. However, the proportion of quenching rates of the present experiments retature-quenched melts (hydrous glass) with in the present paper as well as elsewhere in the complexes. Before discussing possible detailed sanous OH complexes (including molecular lierature, the spectra were obtained from temwhillity mechanisms, it should be noted that leular H<sub>2</sub>O and in the form of various OH 8.0 is consistent with water dissolved as moquenched melts on the join CaALO,-SiO,the interpretation of the Raman spectra of

Stolper and coworkers in several reports 1982; Stolper, 1982; Silver and Stolper, 1989) suggested that the water speciation is only seakly temperature dependent. On the other hand, Dingwell and Webb (1989a, b) suggested that speciation in silicate metrs in general and OH /H<sub>2</sub>O in particular, was signifficantly temperature-dependent. Dingwell and webb (1989b) calculated from relaxation theway, for example, that for a reaction:

20H (\*H,O+O)

that could describe the equilibrium between hetween oxygen in the melt and the two principal forms of water, AH = 25 + 5 kJ mol<sup>-1</sup>. They concluded that OH /H.O measured from quenched materials is that frozen in near the glass transition temperature, and that at the temperature of equilibration, nearly all the dissolved water exists in the OH form. In contrast, Silver and Stolper (1989) suggested that reaction (3) was only weakly temperature-dependent. Unfortunately, until in situ, high-temperature, high-pressure spectroscopy of hydrous melts is carried out, this controversy will remain unresolved, and measurements of OH /H.O of quenched melts should be viewed with these possible problems in mind.

signments. Therefore, the existence of such OH cannot be used with confidence for such ascomplexes in the melts was inferred from the poor, and this portion of the Raman spectra moto, 1978), where the spectral resolution is occur at frequencies ≤600 cm<sup>-1</sup> (e.g., Naka-OH and Al-OH stretch vibrations are likely to of Car OH and AI=OH bonds, the relevant Ca-Carl or with AIV. Due to the ionic character nisms. Either OH groups are associated with CaAl<sub>2</sub>O<sub>4</sub>-SiO<sub>2</sub>-H<sub>2</sub>O quenched melts could be bridging oxygen could from via two mechanonbridging oxygens have been formed by soassigned to (Si,Al)-O stretching. That is, 900 cm 1 for certain compositions) in the ever, the band near 970 cm., and that near lution of H<sub>2</sub>O in CaAl<sub>2</sub>O<sub>4</sub>-SiO<sub>2</sub> melts. Nonmeits in the system NaAlO,-SiO,-H.O. How-(1986) and Mysen and Virgo (1986a) for also reached by McMillan and Remmete groups bonded to Si41 are not of major impor-Oll | stretching leads to the suggestion that Oll that could be assigned to Si-OH [or (Si,At)pleves. The absence of a clear 970-cm 1 band existence of several different M\_OH comsition quenched melts are consistent with the tauce in aluminosilicate metts -- a conclusion The spectra of CaALO,-SiO,-H.O compo-

this discussion above. lopology of the 3000–4000-cm<sup>-1</sup> envelope in

envelope yields direct information about the actions that relate the formation of each of solved water depolymerizes the melt. One can plexing), the reaction is: For the formation of Ca(OH)<sub>2</sub> (Ca..OH comtion of nonbridging oxygen in the melt (NBO). these OH complexes separately to the formawrite, for example, idealized end-member rethe extent to which a fixed amount of disproportion of the various complexes governs plexing with OH in the melts. In principle, the relative importance of Ca2+ and Al3+ comnor that of the bands in the 3000-4000-cm-1 Neither the discussion of the 970-cm<sup>-1</sup> band

$$Ca_nAl_{2n}Si_{2m}O_{4(n+m)} + nH_2O = nCa(OH)_2 + 2nAl^{3+} + 3nSiO_3^2$$

$$+(2m-3n)SiO_2$$
 (4)

$$Ca_{n}Al_{2n}Si_{2m}O_{4(n+m)} + 3nH_{2}O^{-1}$$

$$2n\text{Al}(\text{OH})_3 + n\text{Ca}^{2+} + n\text{SiO}_3^{2-}$$

$$+ (2m-n)SiO2$$
 (5)

analogous to that proposed by Burnham (1974 If water dissolved by replacing a bridging oxy-[(Si,Al)O(OH)], one can write a reaction 1975) for NaAlSi<sub>3</sub>O<sub>8</sub> melt: the aluminosilicate network

$$Ca_nAl_{2n}Si_{2m}O_{a(n+m)} + \frac{3}{2}nH_2O =$$

$$nCa(OH)_2 + Al_{2n}Si_{2m}O_{4m+3n}(OH)_n$$
 (6)

need to charge-balance tetrahedrally-coordiare formed in reaction (6). This effect of dissolved as Al(OH), No nonbridging oxygens NBO will be formed per molecule H₂O dismelt as Ca(OH)2, whereas for reaction (5) be formed per molecule H2O dissolved in the For reaction (4), 6 nonbridging oxygens will (Mysen and Virgo, 1986a), is the result of the for melts in the system NaAlO2-SiO2-H2O solved water on the melt structure, also found

> drous melt become network modifiers. Each of originally in tetrahedral coordination in anhyaction between Al3+ and H2O. This reaction ygens. Thus, for each Ca(OH)2 formed by this ral network can stabilize three nonbridging oxthese Al3+ cations expelled from the tetrahedacts with H<sub>2</sub>O to form Ca(OH)<sub>2</sub>, two Al<sup>1</sup> nated Al3+ with Ca2+. Thus, if one Ca2+ inter formed per molecule H2O dissolved as requires three H2O, and, thus 3NBO will be per two Al(OH)3 complexes formed by internonbridging oxygens will be stabilized the melt fying for two Al3+ forming Al(OH)3. Thus.? formed, one Ca2+ will become network-modi-If, on the other hand, Al(OH)3 complexes are mechanism, 6 nonbridging oxygens are formed

so that the more aluminous the melt by more  $SiO_2/SiO_3^{2-}$ , is a function of n/m (Table VI) cations (NBO/T), does depend on the Al gree of polymerization expressed as nonbride pendent of the Al/(Al+Si). However, the debonded to H in reactions (3) and (4) is indemolecule dissolved H2O as an OH complex not melt depolymerization with increasing water Ca(OH)<sub>2</sub> rather than Al(OH)<sub>3</sub>, the rate of Moreover, by dissolving water in the form of dissolved as either Ca(OH)2 or Al(OH). ing oxygens per tetrahedrally coordinated content is greater. rapidly will the NBO/T increase with water (Al+Si) because the abundance ratio The number of nonbridging oxygens per

In order to evaluate the relative importance

#### TABLE VI

Rate of change of NBO/T with H<sub>2</sub>O concentration in W complexes  $\{\partial(NBO/T)/\partial H_2O\}$ 

0.125 0.200 0.250 0.250 0.333	Al/(Al+Si)	
0	$\frac{\partial (NBO/T)/\partial (H_2O) (mol^{-1})}{(Si_2AI)O(OH)} = AI(OH)_3$	
سانې — اېټ ب⊲ دې سريد ا	2O) (mol <sup>-1</sup> ) Al(OH) <sub>3</sub>	
- ۱۰۰۰ سامع — بیرا	Ca(OIII.	

cens in that unit,  $X_{900}^{NBO/T}$ : dermine the abundance of nonbridging oxy  $\{A_{1150}+A_{1200}\}$ , could then be used to for a given area [e.g.,  $A_{\text{WIR}}^R = A_{\text{WIR}}$ ] Mysen, 1990). Such a calibration factor, a man bands from nonbridging oxygens. These nons (4-6)] it is necessary to determine the urea ratios have been calibrated against NBO/ where the  $A_{900}$  and  $A_{470}$  are the areas of the Ru- $44_{000}+A_{1180}+A_{1200}$ ) and  $A_{970}/(A_{1180}+A_{1200})$ be determined from area ratios such as Annel the melts. In principle, the value of NBO/T can I in the systems Na,O-SiO, and K,O-SiO VBO/T and the abundance ratio, OH/H<sub>2</sub>O, in of the three idealized reaction above [reac

$$V_{\mu\nu}^{\text{VBO/T}} = \theta A_{\text{VBO}}^{R} \tag{7}$$

furthermore, because, with the exception of  $\approx 10$ -wt.% H<sub>2</sub>O isochore, the  $A_{van}/$ procentration range studied. This conclusion for SiO<sub>3</sub><sup>2</sup> units from Mysen (1990) is appliaccording to reaction (6), or (2) the addirelis with > 3 wt.%  $\rm{H}_2O$  is dissolved either mal water is dissolved in molecular form. ronly reasonable if: (1) either the H2O in  $\alpha$  Table V is independent of  $H_2O$  in the water wild be in the range between  $\sim 0.25$  and  $\mathfrak{A}$  .). If it is assumed, nevertheless, that the  $\theta$ factions (i.e. inferred values of  $X_{\text{with}}^{\text{NIOM}}$  from  $\cdot 0.5$ . By assuming that  $\theta$  is independent of he NBO/T-values for the melts studied here VI+Si) and the  $A_{970}/(A_{970}+A_{1180}+A_{1200})$  $t_{ca}+A_{1150}+A_{1200}$ ) decreases with A1/ able to both the  $\nu_{900}$  and  $\nu_{970}$  bands ( $\theta = 1.4$ ). Where systems would underestimate the mole/ "ee melts, and the calibration factor from the re likely to be smaller than in equivalent Alnlensity ratios for a given balk melt NBO/1 ands increase with increasing Al/(Al+Si). (481) that the intensity of the  $\nu_{1180}$  and  $\nu_{1200}$ whatituted for Si4+ in the structural units of omposition, the NBO/T from the area data thus, in the spectra reported here, the relevant sterest. It is known (Sharma and Simons, alibration factors may be affected by AI'+ However, it is not well established how these

> solved H<sub>2</sub>O depolymerize the melt. This trend content of the melt. is more pronounced the lower the total water coordinated Al3+, the less efficiently will dismelts with Ca-charge-balanced tetrahedrally-Thus, the more aluminous aluminositicate plexes and bulk melt Al/(Al+Si)] exists. tween Al(OH<sub>3</sub>/Ca(OH)<sub>2</sub> in the OH comanalogous relation [positive correlation beto a positive correlation between Al/(Al+Si) data that in the system CaAl<sub>2</sub>O<sub>4</sub>-SiO<sub>2</sub>-H<sub>2</sub>O an and Al(OH)3/NaOH in the OH complexes in also observed in the system NaAlO<sub>2</sub>-SiO<sub>2</sub>-H<sub>2</sub>O come more aluminous. The latter trend was the melts. It would appear from the present (Mysen and Virgo, 1986a), and was ascribed water content would decrease as the melts beonly increases slowly with Al/(Al+Si) (see Table V), the NBO/T of a melt for a giver

uminous. This trend results in less efficient de-OH (or Na-OH) as the melts become more alpolymerization of the metts the more alumi bonds become more important relative to Caand Ca-aluminosilicate melts, the Al-OH melts is relatively unimportant. In both Naabundance relative to other OH bonding in the However, if such bonds are present, their AlO<sub>2</sub>-SiO<sub>2</sub> system (Mysen and Virgo, 1986a). scopic observations from melts in the Nauminosilicate melts. In contrast to the spectrotous they are. (Si,Al)-OH bonding cannot be ruled out. predominantly Na..OH complexing in Na-alin the Ca-aluminosificate melts similar to the but probably with Ca complexing dominating with all the cations present (including H+). aluminoslicate melts to form OH complexes In summary, water is dissolved in calcium

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