



Reprint Number 3 3 AD AO 61 495 NEGATIVE-U STATES IN THE GAP IN HYDROGENATED AMORPHOUS SILICON Ronald/Fisch and Donald C./Licciardello Department of Physics Princeton University Princeton, N.J. 08540 NOV 21 1978 Phys. Rev. Lett. 41, 889 (1978) Preprint (Reprint not yet available)/ al Report Date: October 1978 Approved for public release; distribution unlimited 111 Prepared for OFFICE OF NAVAL RESEARCH (Code 427) Arlington, Va 22217 Under Contract N00014-77-C-0711, NR 318-058 NSF-DMR78-\$3\$13 Monitoring Office 42 Office of Naval Research (Code 471) Arlington, Va. 22217

Reproduction in whole or in part is permitted for any purpose of the United States Government

402 527

11 15 020

	ATION OF THIS PAGE (When Data Entered)	
REI	PORT DOCUMENTATION PAGE	READ INSTRUCTIONS BEFORE COMPLETING FORM
I. REPORT NUMBER	2. GOVT ACCESSION NO	2. 3. RECIPIENT'S CATALOG NUMBER
4. TITLE (and Subtitle	e)	S. TYPE OF REPORT & PERIOD COVERE
NEGATIVE-	U STATES IN THE GAP IN HYDROGENATED	Reprint
		6. PERFORMING ORG. REPORT NUMBER
7. AUTHOR(.)		8. CONTRACT OR GRANT NUMBER(*)
R. Fisch a	and D.C. Licciardello	N00014-77-C-07114
9. PERFORMING ORG	ANIZATION NAME AND ADDRESS	10. PROGRAM ELEMENT, PROJECT, TASK AREA & WORK UNIT NUMBERS
	Department	AREA & WORK UNIT NUMBERS
Princeto	on University	NR 318-058
	DD. N.J. 08540	12. REPORT DATE
Office of Arlington,	Naval Research (Code $\frac{42.7}{471}$)	13. NUMBER OF PAGES
	, Va. 22217 INCY NAME & ADDRESS(II dillerent from Controlling Office)	15. SECURITY CLASS. (of this report)
		154. DECLASSIFICATION/DOWNGRADING SCHEDULE
	ATEMENT (of this Report) For public release; distribution unli	mited
Approved f		
Approved f	For public release; distribution unli	
Approved f 17. distribution st 18. supplementary Published	Tor public release; distribution unli ATEMENT (of the abstract entered in Block 20, 11 different f	rom Report)
Approved f 17. DISTRIBUTION ST 18. SUPPLEMENTARY Published 19. KEY WORDS (Contr 19. KEY WORDS (Contr	Tor public release; distribution unli ATEMENT (of the abetract entered in Block 20, 11 different f r NOTES : Phys. Rev. Lett. <u>41</u> (1978) 889	rom Report)
Approved f 17. DISTRIBUTION ST 18. SUPPLEMENTARY Published 19. KEY WORDS (Conti Amorphou Amorphou	Tor public release; distribution unli ATEMENT (of the abstract entered in Block 20, 11 different f r NOTES Phys. Rev. Lett. <u>41</u> (1978) 889 inue on reverse elde if necessary and (dentify by block number is Semiconductors is Silicon	rom Report)
Approved f 17. DISTRIBUTION ST 18. SUPPLEMENTARY Published 19. KEY WORDS (Conti Amorphou Amorphou	Tor public release; distribution unli ATEMENT (of the ebstrect entered in Block 20, if different f r NOTES : Phys. Rev. Lett. <u>41</u> (1978) 889 inue on reverse elde if necessary and (dentify by block number is Semiconductors	rom Report)
Approved f 17. DISTRIBUTION ST 18. SUPPLEMENTARY Published 19. KEY WORDS (Conti Amorphou Amorphou Negative	Tor public release; distribution unli ATEMENT (of the abstract entered in Block 20, if different is r NOTES Phys. Rev. Lett. <u>41</u> (1978) 889 inue on reverse eide if necessary and (dentify by block number s Semiconductors s Silicon -U States	rom Report)
Approved f 17. DISTRIBUTION ST 18. SUPPLEMENTARY Published 19. KEY WORDS (Conti Amorphou Amorphou Negative 20. ABSTRACT (Contin We prop amorphous sili which have a r	Tor public release; distribution unli ATEMENT (of the abstract entered in Block 20, 11 different f NOTES Phys. Rev. Lett. <u>41</u> (1978) 889 Inue on reverse elde 11 necessary and identify by block number as Semiconductors s Silicon -U States Note on reverse elde 11 necessary and identify by block number pose that Si-H-Si three center bonds icon. These centers give rise to state negative effective electronic correlation and the known properties of this to	"" "" " e exist in hydrogenated ates in the energy gap ation energy, U. Our model material. We make suggestion
Approved f 17. DISTRIBUTION ST 18. SUPPLEMENTARY Published 19. KEY WORDS (Conti Amorphou Amorphou Negative 20. ABSTRACT (Contin We prop amorphous sili which have a r	Tor public release; distribution unli ATEMENT (of the abstract entered in Block 20, if different is r NOTES Phys. Rev. Lett. <u>41</u> (1978) 889 inue on reverse elde if necessary and (dentify by block number as Semiconductors s Silicon -U States nue on reverse elde if necessary and identify by block number pose that Si-H-Si three center bonds icon. These centers give rise to state negative effective electronic correlation any of the known properties of this to obtain materials which may prove use	"" "" " e exist in hydrogenated ates in the energy gap ation energy, U. Our model material. We make suggestion
Approved f 17. DISTRIBUTION ST 18. SUPPLEMENTARY Published 19. KEY WORDS (Conti Amorphou Amorphou Negative 20. ABSTRACT (Contin We prop amorphous sili which have a r can explain ma about how to	Tor public release; distribution unli ATEMENT (of the abstract entered in Block 20, if different is r NOTES Phys. Rev. Lett. <u>41</u> (1978) 889 inue on reverse elde if necessary and (dentify by block number as Semiconductors s Silicon -U States nue on reverse elde if necessary and identify by block number pose that Si-H-Si three center bonds icon. These centers give rise to state negative effective electronic correlation any of the known properties of this to obtain materials which may prove use	"" "" " e exist in hydrogenated ates in the energy gap ation energy, U. Our model material. We make suggestion

Submitted to Physical Review Letters 41, 889(1)7.

Negative U States in the Gap in

Hydrogenated Amorphous Silicon*

R. Fisch

and D.C. Licciardello

Department of Physics Princeton University Princeton, New Jersey 08540

USTIFICATION PY DISTRUCTION (AVAILABILITY D ACL GOV / OF ST	TIS DC NANNOUNCED	White Set Buff Séci	an a
DISTRICT ON AMAILABILITY D	USTIFICATION		
A	DISTRICT ON	WAILABILITY and / or	51 51

ABSTRACT

We propose that Si-H-Si three center bonds exist in hydrogenated amorphous silicon. These centers give rise to states in the energy gap which have a negative effective electronic correlation energy, U. Our model can explain many of the known properties of this material. We make suggestions about how to obtain materials which may prove useful in electronic device applications.

*Work supported in part by NSF DMR 78-03015 and ONR N00014-77-C-0711.

PACS Nos. 72.80.Ng, 71.25.Mg

Anderson proposed a model¹ for the electronic Several years ago structure of amorphous semiconductors. The key feature of this model was that the effective correlation energy, U , between two electrons in the same state was hypothesized to be negative, so that the ground state of the material was diamagnetic. Subsequently, Mott, Davis and Street.² and Kastner, Adler and Fritzsche³ identified specific chemical configurations which occur in lone-pair semiconductors and which are responsible for the existence of negative U states in the energy gaps of these materials. These authors claimed that tetrahedrally bonded semiconductors (Si,Ge) cannot form such negative U centers, and thus are fundamentally different from the lone-pair semiconductors. However, Knights, Biegelsen and Solomon⁴ have performed ESR experiments which indictate that negative U centers do exist in hydrogenated amorphous silicon. In this Letter, we point out that the incorporation of large amounts of hydrogen into amorphous Si (or Ge) will allow the formation of a specific type of negative U gap state. A previous attempt to explain the properties of this material assumed the existence of a different type of negative U center. We briefly discuss the implications of our work for the device applications of amorphous silicon-based materials.

The fundamental idea is the following. Due to the random nature of the material, we expect that the Si-Si bonds will have a distribution of energies. Most of the weak bonds will have a larger than normal Si-Si bond length. If there is enough room in the local environment, we will be able to insert two H atoms, break the Si-Si bond, and form two Si-H bonds. If there is somewhat less room available, we will only be able to insert one H atom, and we will form an Si-H-Si three-center bond (TCB). This possibility has been suggested by Mott, Davis and Street. Appelbaum,

-2-

Hamann, and Tasso⁶ have discussed a chemisorptive bond of this type which may occur on the surface of crystalline Si . If we assume for the moment that the H sits halfway between the two Si's, we can form the following molecular orbital wave functions from the atomic orbitals ϕ_{Si_1} , ϕ_{Si_2} , and ϕ_{H} :

$$\psi_{1} = (2+\alpha^{2})^{-\frac{1}{2}}(\phi_{\text{Si}_{1}} + \alpha\phi_{\text{H}} + \phi_{\text{Si}_{2}})$$

$$\psi_{2} = 2^{-\frac{1}{2}}(\phi_{\text{Si}_{1}} - \phi_{\text{Si}_{2}})$$

$$\psi_{3} = (2+4/\alpha^{2})^{-\frac{1}{2}}(\phi_{\text{Si}_{1}} - (2/\alpha)\phi_{\text{H}} + \phi_{\text{Si}_{2}}) .$$
(1)

The wavefunctions ψ_i will, of course, change if we vary the position of the proton. For example, if we move the H away from the center position toward one of the Si's, the ψ_1 orbital will be well approximated by an Si-H bonding orbital. Under these conditions the ψ_2 orbital is approximately an Si lone pair orbital, and the ψ_3 becomes the Si-H antibonding orbital.

 ψ_1 has an energy which is deep in the valence band, and thus will always be occupied by two electrons. The ψ_1 orbital provides the binding energy which keeps the H atom in position between the two Si's. If only the ψ_1 orbital is occupied, the H atom should indeed be stable at a position equally distant from the two Si atoms (although <u>not</u> necessarily on the line between them) since this minimizes the electronic energy. We shall call TCB's which are occupied by 2,3, and 4 electrons T^+ , T^0 , and T^- states, respectively. Note that the superscript corresponds to the local charge of the bond.

The T^+ state is very similar to the TCB's which occur in diborane, B₂H₆. Based on the known structure of the diborane, in which the B-H bond length is 0.14 Å bigger in the TCB configuration than in the normal B-H

-3-

bond, we estimate that the Si-H bond length in this T^+ state is about 1.6 Å. (Recall that the normal Si-H bond is 1.48 Å.) The bond energy should be relatively insensitive to the Si-H-Si bond angle, since ϕ_H is an s orbital. This implies that the bond energy may be approximately constant over a range of Si-Si distances.

Now let us consider the case of the TCB occupied by four electrons. If we fill up the ψ_1 and ψ_2 orbitals, we see that, due to the node in the ψ_2 wavefunction, the center of the bond is no longer a stable The H sees a double-bottomed potential position for the H atom. well, with a maximum in the center. Thus the H will move close to one of the Si's, into the "hydrogen bond" configuration. Because of the local charge, the T state forms a "strong hydrogen bond", 7 which lowers its energy relative to the T⁰ state. Thus the ψ_2 orbital has a negative U whose magnitude can be estimated to be 0.6-1.0 eV. distance over which the H moves on going from the T state to the T state should be at least 0.15-0.20 Å , and it may be considerably larger, depending on the change in the Si-Si distance. This latter quantity is difficult to estimate, since it depends heavily on the local environment surrounding the bond. We believe that most of the motion of the proton occurs upon addition of the first electron to the ψ_2 orbital.

Since ψ_2 is essentially a "non-bonding" orbital, we anticipate that it will have an energy which lies in the gap between the valence and conduction bands. (Note that we must not use the wavefunction of Equation (1) to calculate the energy of ψ_2 , since when ψ_2 is occupied the H is not in the centered position.) Because of the negative U, in the ground state we must put electrons into the ψ_2 orbitals in pairs. By charge neutrality, we see that in undoped a-Si we can fill half of the available ψ_2 orbitals. Since all spins are paired, the ground state of a TCB will be diamagnetic.

-4-

The TCB's can be considered to be canonical Anderson-Halperin-Varma^o tunneling centers. They give linear contributions to the low-temperature specific heat in two ways. First, there is an electronic contribution which results from the movement of pairs of electrons among the available orbitals. Second, there is a "phonon" contribution which results from the hopping of "hydrogen bond" protons from one side to the other of their double wells. (Due to local environment effects, the two sides will not be precisely equivalent.) It is not clear that the number of TCB's (not more than $10^{18}/cm^3$) is large enough to explain the magnitude of the linear specific in this regard heat. Their importance can easily be checked by making measurements on lightly-doped samples.

The T^+ and T^- states constitute a lower Hubbard (LH) band, with the T^0 states making up the upper Hubbard (UH) band. (Recall that U is negative.) From experimental measurements,⁹ we can place the LH band at about 0.6 eV below the conduction band. Our estimate of U places the UH band just below the conduction band, since the UH band lies |U/2| above the LH band. This density of states is shown schematically in Figure 1. It is important to note that one cannot simply obtain the density of gap states as a function of energy, n(E), by varying the density of electrons in the material. This is because the correlation energy, U, must be accounted for, since the LH band consists of twoparticle states rather than single-particle states.

From experiment,⁹ we know that the energy gap between the valence and conduction band mobility edges $Eg \cong 1.7 \text{ eV}$. Thus, if we define W to be the energy difference between the valence band mobility edge and the

-5-

Fermi energy in the undoped material, $W \cong 1.1 \text{ eV}$. We can estimate the energies of the various electronic transitions which can occur in terms of these quantities:

electron + hole
$$\neq$$
 hv $\Delta E = Eg \approx 1.7 \text{ eV}$ (2)

electron +
$$T^{+} \stackrel{\downarrow}{\leftarrow} T^{0}$$
 $\Delta E = Eg - W - \frac{|U|}{2} \approx 0.2 \text{ eV}$ (3)

hole +
$$T^{-} \stackrel{?}{\leftarrow} T^{0}$$
 $\Delta E = W - \frac{|U|}{2} \approx 0.7 \text{ eV}$ (4)

electron +
$$T^{0+}_{+}$$
 T $\Delta E = Eg - W + \frac{|U|}{2} = 1.0 \text{ eV}$ (5)

hole +
$$T^0 \stackrel{?}{\leftarrow} T^+$$
 $\Delta E = W + \frac{|U|}{2} \approx 1.5 \text{ eV}$ (6)

Equation (2) represents the intrinsic band edge recombination process. Equations (3) and (4) represent the trapping of electrons and holes, respectively. We see that T⁻ centers act as deep hole traps, but T⁺ centers are shallow electron traps. This is in excellent agreement with the experimental result¹⁰ that the electron mobility is more than an order of magnitude greater than the hole mobility. Equations (5) and (6) represent the creation of T⁻ and T⁺ centers, respectively. All of these processes should give rise to photoluminescence. The creation of T⁺ centers, Equation (6), should have an appreciable Stokes shift, because of the associated motion of the H atom. Our model's predictions agree very well with the experimentally observed photoluminescence.¹¹⁻¹⁴

Since amorphous silicon is a complex material, it would be unreasonable to expect that all states in the gap arise from the same mechanism. Residual dangling bond states and band tail states undoubtedly do exist, but these types of states should have a positive U.

-6-

Our model provides a basis for making suggestions of what one might do to obtain technologically useful material. At this point, the major limitation in the use of hydrogenated a-Si for solar cell applications is the poor hole mobility¹⁰ of the n-type material. Our model predicts that if we make lightly boron-doped (about 50 ppm) material, we will remove most of the T⁻ states. Thus, we may obtain material which has a very small number of thermally excited carriers, but also has a relatively high carrier mobility.

Whether this type of material will work efficiently in solar cells depends on the rate at which T⁻ centers are created by recombination processes, Equation (5), and whether there exist other types of hole traps.

Alternatively, we may attempt to remove the TCB's from the material altogether. This, of course, requires removing all of the hydrogen from the material. As a substitute for hydrogen, we may consider helogenated amorphous silicon. Halogen atoms act to tie up all of the dangling Si bonds, but they cannot form TCB's as hydrogen can. We suggest that fluorine is the most likely candidate for obtaining useful material for two reasons. First, the length of the Si-F bond (1.56 Å) is only slightly greater than the length of the Si-H bond (1.48 Å), while Si-Cl (2.03 Å) and Si-Br (2.17 Å) are significantly larger. Second, SiF₄, BF₃, and PF₃ are all gases at room temperature, which may prove convenient for materials preparation.

In summary, we have proposed that hydrogenated amorphous silicon contains significant numbers of Si-H-Si three-center bonds. Since these centers have states which lie in the energy gap and have a negative

-7-

can effective electronic correlation energy U, their existence explain much of the heretofore puzzling experimental information about these materials. Our model allows us to make predictions about how to obtain technologically more useful amorphous silicon-based materials.

Acknowledgements

We thank Professor L.C. Allen and Dr. D.R. Hamann for helpful discussions, and Mr. J.J. Park for bringing Reference 5 to our attention. We are grateful to Professor J.J. Hopfield for his advice and encouragement.

-8-

References

1.	P.W.	Anderson,	Phys	Rev.	Lett.	34.	953	(1975).	

- N.F. Mott, E.A. Davis, and .R.A. Street, Phil. Mag. <u>32</u>, 961 (1975);
 R.A. Street and N.F. Mott, Phys. Rev. Lett. 35, 1293 (1975).
- 3. M. Kastner, D. Adler, and H. Fritzsche, Phys. Rev. Lett. 37, 1504 (1976).
- J.C. Knights, D.K. Biegelsen, and I. Solomon, Solid State Commun. <u>22</u>,
 133 (1977); J.C. Knights, <u>Proc. Seventh Intl. Conf. on Amorphous and</u> <u>Liquid Semiconductors</u>, Edinburgh 1977, W.E. Spear, ed. (Univ. Edinburgh, 1977), p. 433.
- D.L. Stein, D.C. Licciardello, and K.B. Ma, Bull. Am. Phys. Soc. <u>23</u>, 414 (1978).
- 6. J.A. Appelbaum, D.R. Hamann, and K.H. Tasso, Phys. Rev. Lett. 39,1487(1977).
- L.C. Allen, J. Amer. Chem. Soc. <u>97</u> 6921 (1975); and private communication.
- P.W. Anderson, B.I. Halperin, and C.M. Varma, Phil. Mag. <u>25</u>, 1 (1972).
 W.E. Spear and P.G. LeComber, Phil. Mag. 33, 935 (1976).
- 10. D.E. Carlson, Amorphous-Silicon Solar Cells, RCA report, Sept. 1977.
- T.S. Nashashibi, I.G. Austin, and T.M. Searle, Phil. Mag. <u>35</u>, 831 (1977).
- 12. D. Engemann and R. Fischer, Phys. Stat. Sol. (b) 79, 195 (1977).
- J.I. Pankove and D.E. Carlson, Appl. Phys. Lett. <u>31</u>, 450 (1977);
 <u>Proc. Seventh Intl. Conf. on Amorphous and Liquid Semiconductors</u>,
 Edinburgh 1977, W.E. Spear, ed. (Univ. Edinburgh, 1977), p. 433.
- 14. R.A. Street, Phil. Mag. 37, 35 (1978).



