Electronic excitations in finite and infinite polyenes

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We study electronic excitations in long polyenes, i.e., in one-dimensional strongly correlated electron systems which are neither infinite nor small. The excitations are described within Hubbard and Pariser-Parr-Pople (PPP) models by means of a multiple-reference double-excitation expansion [P. Tavan and K. Schulten, J. Chem. Phys. 85, 6602 (1986)]. We find that quantized "transition" momenta can be assigned to electronic excitations in finite chains. These momenta link excitation energies of finite chains to dispersion relations of infinite chains, i.e., they bridge the gap between finite and infinite systems. A key result is the following: Excitation energies \( E \) in polyenes with \( N \) carbon atoms are described very accurately by the formula \( E = \Delta E_0 + \alpha^q (N)q \), \( q = 1, 2, \ldots \), where \( \Delta E_0 \) denotes the excitation class, \( \Delta E_0 \) the energy gap in the infinite system \( [\alpha^0 (N) > 0] \), and \( k (N) \) the elementary transition momentum. The parameters \( \Delta E_0 \) and \( \alpha^q \) are determined for covalent and ionic excitations in alternating and nonalternating polyenes. The covalent excitations are combinations of triplet excitations \( T \), i.e., \( T, TT, TTT, \ldots \). The lowest singlet excitations in the infinite polyene, e.g., in polyacetylene or polydiacetylene, are \( TT \) states. Available evidence proves that these states can dissociate into separate triplets. The bond structure of \( TT \) states is that of a neutral soliton-antisoliton pair. The level density of \( TT \) states in long polyenes is high enough to allow dissociation into separate solitons.

I. INTRODUCTION

Motivated by the aim to develop conducting and semiconducting organic materials the properties of conjugated polymers, in particular, the properties of their prototype \textit{trans}-polyacetylene \([\text{trans-}(CH)_3]\) (see Fig. 1), have been investigated in recent years.\(^1\) The investigations have been guided by solid-state-physics concepts developed for "infinite"-size materials, i.e., materials characterized by bands of very densely spaced electronic states. However, the organic conductors and semiconductors investigated so far are by no means infinite. Polyacetylene, for instance, consists of strands of conjugated polymers, called polyenes, which are some hundred carbon atoms long. It is not known how densely spaced the electronic excitations in such chains actually are and to what extent a mixing of electronic states is possible. In this paper we will attempt to bridge the gap between finite and infinite polyenes. The results will be rewarding: an extrapolation of the well-known properties of small polyenes\(^2\) to very long polyenes provides an estimate of the density of states of finite, but long polyenes, yields explanations for previous observations, suggests new observations, and sheds light on the concepts employed so far to understand the properties of semiconducting organic polymers.

The concept that has been envoled most often in the field of organic polymers in the recent past is that of the soliton, i.e., of a low-energy excitation that involves a strong coupling between electronic and lattice degrees of freedom. It has been suggested that solitonic excitations mediate the conductivity of \((CH)_x\). In the pioneering work of Su, Schrieffer and Heeger the electron-phonon coupling had been emphasized at the expense of electron-electron interactions which had been neglected.\(^3\)\(^4\) Subsequent theoretical studies on conjugated polymers have attempted, therefore, to evaluate the effect of electron-electron interactions on the energies and lattice structures of solitonic excitations in electron-phonon models of polyacetylene.\(^5\)\(^9\)\(^11\)

Solitonic excitations as suggested by Su, Schrieffer, and Heeger have not been observed in small polyenes. What has been observed are vibronic effects on isolated electronic states, e.g., shifts of equilibrium geometries and vibrational frequencies upon excitations from the ground state to isolated excited states. The question arises whether the properties of long polyacetylene strands \((CH)_x\) arise from isolated stationary, delocalized states as in shorter polyenes or whether wave packets combined from nearly degenerate delocalized electronic excitations and shaped by a coupling to lattice distortions manifest themselves. Solitons belong to the latter class of excitations. For an answer to the question raised one needs to extrapolate the properties of finite polyenes to the long polyene limit in order (1) to see how the isolated electronic excitations in

FIG. 1. Ground state of polyenes and polyacetylene characterized by alternating single and double bonds.
the short polyenes merge to form bands of excitations in the longer polyenes, (2) to obtain information on the properties of the respective excitations, e.g., their bond pattern, and (3) to estimate the intraband state density of long, but finite polyenes. This latter information determines to what extent weak interactions like electron-phonon coupling can induce new states, e.g., solitonic excitations in \((\text{CH})_x\).

The following results of work on finite polyenes appear to be relevant for current investigations on polyacetylene.

(1) Qualitatively correct and quantitatively accurate descriptions of electronic spectra require highly sophisticated many-electron methods because the motion of electrons in these one-dimensional materials is dominated by Coulomb repulsion.

(2) The size of the energy gap between the ground state and the first optically allowed excited singlet state is determined by the strength of the Coulomb interaction between the \(\pi\) electrons; bond alternation, i.e., the pattern of single (long) and double (short) bonds, contributes only a minor fraction to the energy gap.\(^{12,13}\) The first optically allowed singlet state is the lowest energy member of a class of ionic singlet states in polyenes which are either optically allowed or assume rather strong oscillator strengths upon a breaking of the molecular symmetry, e.g., upon trans \(-\) cis isomerization.\(^2\)

(3) In polyenes there exist optically forbidden covalent singlet electronic excitations with energies below the "optical" gap. These states are characterized by collective excitations of the spin degrees of freedom of the \(\pi\) electrons and, therefore, do not emerge in single-electron theories as these theories largely neglect the effect of electron correlation.\(^{14,15}\) The excitation energy of the covalent states depends sensitively on the degree of bond alternation, i.e., calculations of the energies of these collective \(\pi\)-electron excitations have to be based on realistic molecular geometries in order to render quantitative descriptions.\(^{13}\) The optically forbidden character of the covalent states is not affected very much by a breaking of the molecular symmetry.

(4) The Born-Oppenheimer approximation applies to calculations of electronic wave functions. Even if pure \(\pi\)-electron Hamiltonians are employed for the description of polyenes the equilibrium molecular geometries in low-energy states can be estimated from calculated wave functions. These wave functions, on the other hand, are rather insensitive to the model geometry chosen for the carbon lattice, i.e., in both ground and excited states the electronic structure determines that of the lattice whereas the reverse influence is smaller. For instance, the well-known ground-state structure of alternating single and double bonds shown in Fig. 1, is revealed by bond orders that result from a calculation which assumes equal lengths for all polyene bonds (see also Sec. VI of this paper). Furthermore, it has been shown that excited state bond orders explain the known photochemical behavior of polyynes.\(^{16}\) Hence, for a description of finite polyenes an inclusion of electron-phonon interaction represents an unnecessary complication.

We note that the ionic states are often termed "charge-transfer" excitations and that the covalent states are often referred to as "spin-wave" excitations.

Unfortunately, the knowledge on finite polyenes had been limited until recently\(^{13}\) to short compounds with up to 10 carbons.\(^{12,15}\) These compounds are too short to identify the asymptotic laws which allow an extrapolation to very long polyenes. The results obtained in Ref. 13 and in this paper alter this situation. We have presented in Ref. 13 accurate descriptions of electronic excitations for polyenes up to 16 carbons long and, in part by using results presented already there, we will reveal in this paper the asymptotic relationships that link short and very long polyenes.

The main result of previous studies on finite polyenes is that electron correlation contributes in an essential way to the properties of these compounds. Recently, the important role of electron correlation has also been deduced by observations on polyacetylene \((\text{CH})_x\) and related materials. The occurrence of negative spin densities on alternate carbon atoms which arises due to electron-electron interaction has been observed.\(^{17}\) Single electron theories, in contrast to more complete theories, predict an absorption due to the neutral soliton at the "midgap" position, i.e., at a position where it has not been found.\(^{18}\) Furthermore, the dependence of photocarriers on the photon energy indicates electron-hole interactions which are also not accounted for by single electron theories.\(^{19}\) Consequently, the important effect of electron correlation on the physical properties of \((\text{CH})_x\) has been emphasized.\(^{20}\)

Up to now most theoretical investigations of correlation effects in \((\text{CH})_x\) dealt with bond alternation,\(^{21-28}\) with the optical gap,\(^{12,29-31}\) and with solitonic excitations\(^{5-11}\) in \((\text{CH})_x\). These studies have demonstrated that an assessment of correlation effects on these properties of \((\text{CH})_x\) depends on the quality of the approximation employed. Hartree-Fock\(^{12,28}\) and perturbative treatments\(^{22,5}\) have been shown to render qualitatively wrong predictions on the implications of electron correlations for soliton energetics\(^{26,11}\) and bond alternation.\(^{24-28}\) Therefore, we expect that limited configuration interaction \((\text{CI})\) descriptions like those in Ref. 9 strongly underestimate the effect of Coulomb interactions on soliton energetics. Any satisfactory description of electron correlation effects requires either exact solutions or extend "size-consistent" CI descriptions. Unfortunately, the enormous dimension of the many-electron Hamiltonian for polyenes with 12 and more carbon atoms makes such calculations a computationally extremely difficult task.\(^{12}\) To surmount this task we have developed a new multiple-reference double-excitation configuration-internal interaction method (MRD-CI) which allows very accurate calculations of the low-energy spectra of polyynes comprising up to 16 carbon atoms. A detailed account of this method, a discussion of its accuracy, the results of Pariser-Parr-Pople (PPP) MRD-CI calculations on polynene spectra as well as a discussion of corresponding spectroscopic observations have been presented elsewhere.\(^{12,32}\)

For an extrapolation of the properties of small \((N \leq 16, N = \text{number of carbon atoms})\) polyynes to long polyenes we investigate first the Hubbard model that is the most simple model to describe electron correlation effects in polyenes. For this model exact dispersion relations for the
infinite polyene are known.\textsuperscript{33–37} Also for this model some of the optically allowed (ionic) excitations for finite ring polyenes are known exactly.\textsuperscript{25} Accurate excitations energies for covalent states can be calculated by the MRD-CI method.\textsuperscript{13} We demonstrate first how dispersion relations of infinite polyenes allow to approximately express electronic excitations of finite ring polyenes. This implies that we determine the momenta \( k(N) \) connected with multielectron excitations in ring compounds of size \( N \). We find that these momenta obey simple quantization rules which can be employed to express whole classes of finite polyene electronic excitations by means of infinite polyene dispersion relations.

We then attempt to link infinite Hubbard chain dispersion relations to excitation energies in finite open Hubbard chains of size \( N \). The latter excitation energies are determined by our numerical procedure. We discover, in fact, a simple asymptotic \((N \to \infty)\) behavior for finite Hubbard chains and, thereby, demonstrate that electronic excitations of finite compounds can be extrapolated accurately to the infinite compound. Next, we follow the same procedure to link finite and infinite polyenes as described by the so-called PPP model. This model provides a realistic description of the long-range character of the electron-electron interaction. Fortunately, the results show a close similarity between the asymptotic \((N \to \infty)\) behavior of Hubbard and PPP chains. This finding justifies extrapolation of finite polyene calculations to polycatenylene and polydiacetylene. The following properties are investigated:

1. the structure of the bands of ionic and of covalent electronic excitations,
2. the level density of these bands as a function of chain length,
3. the bond structure of ground state and ionic and covalent excited states.

An important result of our paper is the characterization of a band of covalent states in infinite polyenes with a band edge below that of the conduction band. The expectation\textsuperscript{19} that a band of "homopolar,"\textsuperscript{36} i.e., covalent, excited states should exist in this energy region has been expressed already previously. This expectation originates from observations on smaller polyenes. As mentioned above it had been demonstrated, that the optically allowed polyene \( \sigma^* \) state, which gives rise to the conduction band of (CH)\textsubscript{\(x\)},\textsuperscript{12} is not the lowest excited singlet state in these molecules. Contrary to the predictions of single-electron theories a strongly correlated "covalent" \( A_\sigma \) state\textsuperscript{38} was found to be the lowest singlet excitation and, consequently, to be involved in the photophysics and photochemistry of polyenes.\textsuperscript{2} Spectroscopic observations and a theoretical description revealed that the energy gap between the excited \( B_\sigma \) and the excited \( A_\sigma \) state increases with increasing polyene length,\textsuperscript{15} a result which suggests the expectation of a low-lying covalent band in (CH)\textsubscript{\(x\)}. The \( A_\sigma \) state has been inferred also to explain the long-lived 1.2 eV luminescence observed at low temperatures in (CH)\textsubscript{\(x\)}.\textsuperscript{39,40}

It is the aim of this paper to substantiate the previous speculations on the band of covalent states and to discuss its relation to so-called midgap excitations.

Our paper is structured as follows. In Sec. II we introduce the MRD-CI method, the Hubbard, and PPP model Hamiltonians and describe briefly the interaction parameters assumed in our calculations. In Sec. III we review the characterization of the classes of electronic excitations in small polyenes. In Sec. IV we investigate the asymptotic behavior of excitations in Hubbard rings and Hubbard chains. In this section we provide the conceptual basis of our subsequent discussions. In Sec. V we extend the investigation to polyenes described by the PPP model. This section and the following sections deal with the properties of realistic molecular systems. In Sec. VI we discuss the bond structure of excitations in very long polyenes as described by the PPP model. In Sec. VII we provide explanations of previous observations on polycatenylene and polydiacetylene compounds and add suggestions for new observations which might reveal the character of mid-gap excitations in polycatenylene. Section VIII summarizes the results and states open questions.

### II. METHODS

In this section we will briefly introduce Hamiltonians and computational methods employed in our calculations. The Hamiltonians were chosen to represent the \( \pi \) electrons in the polyenes and polycatenylene, and in particular, the effect of electron correlation. For this purpose the \( \pi \)-electron Hamiltonians must include at least a kinetic energy term that describes the delocalization of electrons, and a Coulomb repulsion term that induces electron correlation. The most simple Hamiltonian of this kind is the Hubbard Hamiltonian

\[
H_{\text{Hub}} = t \sum_{n,m,\sigma} c_{n,\sigma}^\dagger c_{m,\sigma} + U \sum_{n,\sigma} c_{n,\sigma}^\dagger c_{n,\sigma} c_{n,-\sigma}^\dagger c_{n,-\sigma} \tag{1}
\]

which characterizes electron systems by a single parameter \( U/|t| \). The fermion creation (annihilation) operators \( c_{n,\sigma} (c_{n,\sigma}^\dagger) \) in this Hamiltonian position the \( \pi \) electrons in an orthonormal set of atomic orbitals labeled by \( n,m \).

The first term in (1) represents the kinetic energy where the corresponding sum includes only nearest-neighbor sites \( n = m \pm 1 \). The second term describes the repulsion between \( \pi \) electrons, the latter interaction being restricted to situations when two \( \pi \) electrons reside at the same atomic site \( n \). The Hubbard Hamiltonian had been considered for polyenes also in Ref. 2 and by Soos and Ramasesha.\textsuperscript{12}

The effective Hamiltonian most often employed for the description of electronic excitations in polyenes is that originally suggested by Pariser, Parr, and Pople (PPP).\textsuperscript{41} This Hamiltonian assumes that the motion of \( \sigma \) and \( \pi \) electrons is separable and accounts only for the \( \pi \) electrons. We have employed the PPP Hamiltonian previously to describe the spectra of polyenes with up to 10 carbons.\textsuperscript{2,14,15,38} The results were in good agreement with spectral observations on these polyenes. We have chosen, therefore, the PPP Hamiltonian also for our work presented in this and a preceding paper\textsuperscript{13} which extend our previous descriptions to long polyenes and polycatenylene. The PPP Hamiltonian employed is in second quantization
The first term describes nuclear repulsion, the second term attraction between $\pi$ electrons and nuclear cores, the third term the kinetic energy of the $\pi$ electrons, and the last term their mutual Coulomb repulsion. The latter is not restricted to an intra-atomic interaction as in Eq. (1) but extends over the whole molecule. The effective Coulomb repulsion integrals between $\pi$ electrons at sites $n$ and $m$ are given by the Ohno formula\(^{42}\)

\[ U_{nm} = \frac{U}{(1 + 0.6117 r_{nm}^2)^{1/2}}. \]  

With a choice of 11.26 eV for the repulsion energy $U$ of two $\pi$ electrons at the same atom site the two-electron matrix elements $U_{nm}$ asymptotically, i.e., at long distances $r_{nm}$, approach the Coulomb interaction $U(r_{nm}) = e^2/r_{nm}$, whereas at short distances the interelectronic forces correspond to those described by the Hubbard model with $U/|t| = 1.53$.

In order to study the effect of a possible screening of the $\pi$-electron Coulomb repulsion due to the presence of electrons in the $\sigma$ core\(^{6-8}\) we have assumed in some of our calculations a short-range Coulomb interaction

\[ U^{\text{sr}}(r) = U(2r), \]  

which we actually described by scaling the distances $r_{nm}$ in (3) by a factor of 2. At large distances this choice implies a dielectric screening like that in a medium with an optical density $n^2 = 2$. At short distances the choice entails increased interelectronic forces corresponding to those described by the $U/|t| = 1.95$ Hubbard model.

Polymers and polyacetylene assume a bond structure that features an alternation of double (short) and single (long) bonds. This bond alternation will be described in our calculations by a parameter $\delta$ which measures the difference of the lengths of single and double bonds. We will consider two models for the polyene geometry, that corresponding to an intermediate degree of bond alternation as observed in rather short polyenes ($N \approx 10$) and as described by $\delta = 0.10$ Å, and that corresponding to equal bond lengths as described by $\delta = 0$.

The bond length affects the resonance integral $t$ in the PPP Hamiltonian (2) that connects neighboring sites. The resonance integral can be expressed in terms of $\delta$ and is given by (distances $\delta$ in Å)

\[ t = -2.4(1.0 \pm 0.76) \text{ eV}, \]  

where the + and − signs refer to double and single bonds, respectively. In our calculations employing the Hubbard model we have considered only polyenes without bond alternation, i.e., with $\delta = 0$.

The parameterization of the PPP Hamiltonian characterized above is identical to that employed by Soos et al. in their exact valence bond calculations on small ($N \leq 12$) polyenes.\(^{12,43,44}\) The interaction parameters deviate slightly from those in our previous work\(^{2,14,15,38}\) and have been adopted to allow comparisons with the results of these authors.\(^{13}\)

Approximate values for the bond lengths $l_n$, $n = 1, \ldots, N - 1$ between neighboring sites $n$ and $n + 1$ can be predicted because of the $\pi$-bond orders $p_n$. The bond orders, defined separately for each electronic state, are the off-diagonal elements $S_{n,n+1}$ of the one-electron density matrix in the atomic orbital basis. The bond lengths can be evaluated by means of the empirical formula (bond lengths $l_n$ in Å)

\[ l_n = 1.5274 - 0.2105 p_n. \]  

Equation (6) represents the linear interpolation which reproduces the observed single- and double-bond lengths for polyenes with four (butadiene) and six carbons.\(^{43,44}\) The bond orders to be used for this purpose are those from a PPP-MRD-CI\(^{14}\) calculation for a zeroth order polyene geometry described by $\delta = 0.10$ Å.

To simplify the discussion in Sec. VI on polyene geometries in different electronic states we will use the order parameter $\Delta l_n$ defined by

\[ \Delta l_n = (-1)^n (l_n - \bar{t}), \]  

where $\bar{t}$ is the average bond length in the respective state. The order parameter measures the degree of bond alternation in a polyene and allows a simple identification of solitonic excitations.

In order to describe accurately the spectra of the Hamiltonians in Eqs. (1) and (2) as well as the electronic properties of the corresponding electronic states we have applied the multiple-reference double-excitation configuration-interaction (MRD-CI) method presented in some detail in Refs. 13 and 32. The MRD-CI scheme\(^{46,47}\) is characterized by an individual selection of the CI basis for each electronic state. First, the so-called "reference configurations" are determined that describe the main contributions to the respective wave function and define the unperturbed state. Then all those configurations are included into the CI expansion that in second-order perturbation theory can give an energy correction to the unperturbed state. The dimension of the resulting MRD-CI eigenvalue problem depends on the choice of the unperturbed state, on the size of the electron system as well as on the physical nature of the respective state. The MRD-CI basis may comprise up to several $10^6$ spin-adapted, antisymmetrized functions. For all polyene states investigated in this paper the description of the unperturbed states has been chosen identical to that presented in Ref. 13. An exception is the ground state of the PPP model without bond alternation ($\delta = 0$). For this state we have chosen only two reference configurations instead of three as in Ref. 13, resulting in a reduction of the predicted excitation energies by $\approx 0.2$ eV. With this choice we achieved a uniform and consistent description of all polyene states in all models considered here. Among these models only the MRD-CI description of the extremely
strong correlation effects in the $U/|t|=4$ Hubbard model deserves additional consideration.\textsuperscript{48} But for the remaining cases all MRD-CI calculations of polyyne ground states are based on a two reference description, for all ionic excited states one reference has been employed and for all covalent excited states the description has been chosen identical to that of the first excited state presented in Ref. 13.

III. CLASSIFICATION OF POLYNYE EXCITED STATES

In the following discussions we will refer to electronic states of polynes by the symbols $A_g$ and $B_u$ employing the notation of the $C_{2h}$-symmetry point group.\textsuperscript{7} The polyyne states are eigenstates of the total spin operator. A singlet total spin is indicated by the superscript 1 in the symbols $^1A_g$ and $^1B_u$. To denote a triplet state the superscript 1 is replaced by 3. We will consider mainly singlet states because ground state and electronic states reached first after optical excitation and internal conversion are of the singlet variety.

The PPP and Hubbard Hamiltonians give rise to yet another symmetry, the so-called "Pariser alternancy symmetry" or "particle-hole symmetry" (see Refs. 49 and 50). This symmetry classifies states, e.g., the ground state as $^1A_g$, and + states, e.g., the lowest optically allowed state as $^3B_u$. Though only approximate for real molecules the alternancy symmetry is useful for two reasons. First, it allows to reduce the computational effort by taking advantage of the fact that the PPP and Hubbard many-electron Hamiltonians do not mix $^1A_g$, $^1A_g$, $^1B_u$, and $^1B_u$ states. Of course, we have taken advantage of this chance. Second, the alternancy symmetry provides a very simple classification of ionic states which are +, and covalent states which are -. We will, therefore, refer to ionic states of polynes and polyacetylene by an index +, and to covalent states by an index -.

The classification of the electronic states is complete when one numbers the states of each symmetry class as $1, 2, \ldots$ in the order of increasing energy. In this notation the ground state is referred to as $^1A_g$.

Below we will also consider electronic excitations of Hubbard ring molecules with $N=4n+2$, $n=1, 2, \ldots$, carbon atoms. The excitations can be classified according to the $D_{2h}$-symmetry point group. The classes of excitations which we will consider derive from the point group $D_{2h}$. The symmetry class for the many-electron states which involve only $\pi$ electrons are $A_g$, $B_{1u}$, $B_{2u}$, $E_{1u}$, $E_{2g}$. For the definition of these classes see Ref. 51. Pariser's alternancy symmetry applies within the Hubbard model to the ring compounds as well and, hence, the excitations can be further labeled as + and - states. For the singlet states the classifications of Hubbard ring excitations are then $^1B_{1u}$, $^1E_{2g}$, etc.

IV. HUBBARD RINGS AND CHAINS

Before we investigate the PPP model that provides a rather realistic description of polynes we consider the less realistic Hubbard model since for this model exact results for the infinite system as well as for finite systems are available. Comparisons of our MRD-CI results on the low-lying singlet excitations of finite systems with exact results on finite as well as on infinite systems will serve us to test how our MRD-CI results on finite systems can be extrapolated to the infinite polyyne limit, i.e., to polyacetylene (CH)\textsubscript{n}. Our analysis of the size dependence of the low-lying excitations of Hubbard rings and Hubbard chains in this section will reveal features that are remarkably similar to those presented in Sec. V for the PPP model.

A. Dispersion relations of the infinite Hubbard chains and their relationship to electronic excitations in finite chains

The electronic excitations of the infinite Hubbard chain form bands which can be classified by certain quantum numbers which derive from the well-known Bethe ansatz.\textsuperscript{52} A most recent study of the subject is that by Woynarovich\textsuperscript{35-37} which provided analytical expressions for the dispersion relations of the bands of electronic excitations in the infinite Hubbard chain. In the following we will consider three dispersion relations which we determined according to the expressions provided by Woynarovich: (1) the dispersion relation for the band of ionic singlet states denoted by $E^+(k)$, where $k$ is the momentum variable; (2) that for the lower edge of the band of covalent singlet excitations denoted by $E^{-1}(k)$; (3) that for the covalent singlet excitations characterized by four holes in the $\lambda$ distribution\textsuperscript{17} denoted by $E^-(k) = \sum_{i=1}^4 E^{-1}(k_i)$ with $k = \sum_{i=1}^4 k_i$. This latter dispersion relation describes the upper edge of the band of covalent singlet states. All dispersion relations depend, of course, parametrically on the Hubbard parameter $U/|t|$.

The lowest transitions to ionic and covalent states of the infinite Hubbard chain have vanishing momenta $k$. The corresponding excitation energies—the "optical gap" $E^+(0)$ and the "covalent gap" $E^{-1}(0)$—have values $E^+(0) \neq 0$ and $E^{-1}(0) = E^+(0) = 0$, respectively. The question arises if the infinite chain dispersion relations can also provide an approximate description of the lowest excitation energies for finite Hubbard chains. For an answer to this question one has to consider that finite molecules cannot accommodate electronic states with vanishing momenta and that finite size molecules do not conserve momentum upon electronic excitation.

An estimate for the smallest momentum change $k$ involved in an electronic excitation of a finite polyyne can be obtained by considering the $U=0$ case for which analytical expressions for the excitation energies exist. In this case the analytical descriptions of the excitation energies $E(N)$ in finite rings and chains derive from the dispersion relations $E(k)$ of the infinite chain. A straightforward calculation yields the following dispersion relation $E(k)$ for the energetically lowest particle-hole excitation

$$E(k) = 4 |t| \sin(k/2),$$

where $k$ is the momentum change connected with the excitation and where the lattice constant has been chosen to be unity. In finite systems the minimum values of $k$ are
\[ k'(N) = \frac{2\pi}{N} \]  \hspace{1cm} (9)

for rings with \( N = 4n + 2 \) carbon atoms, and

\[ k(c)(N) = \frac{\pi}{N + 1} \]  \hspace{1cm} (10)

for chains with \( N = 2n \) carbon atoms. The analytical expressions for the excitation energies \( E_{k'}^r(N) \) and \( E_c^l(N) \) of the lowest particle-hole excitations in rings and chains, respectively, are now obtained by inserting \( k' \) and \( k_c \) given by Eqs. (9) and (10) into the dispersion relation given by Eq. (8). Similarly, excitation energies \( E_{k'}^c \) and \( E_c^c \) for further transitions are obtained if one inserts multiples of \( k' \) and \( k_c \) into Eq. (8), i.e., for the rings the momenta

\[ k_q' = qk' \]  \hspace{1cm} (11)

and for the chains the momenta

\[ k_q^c = qk_c \]  \hspace{1cm} (12)

where \( q = 1, 2, \ldots \) defines a momentum quantum number characteristic for the respective transition. From now on we will use the term transition momentum for the momentum change involved in an electronic excitation.

For a system of independent electrons the transition momenta are solely determined by the geometry of the molecule. In a correlated electron system the character of an excitation can depend on the strength of the electron-electron repulsion \( U \). For such excitations the transition momenta can vary with \( U \). If the character of the excitation is independent of \( U \), the values of the transition momenta provided by Eqs. (8)–(12) for the \( U = 0 \) case apply to all \( U \) values. This is the case, actually, for finite Hubbard rings for which the total momenta of the charge-transfer as well as of spin-wave excitations are integer (or half-odd integer) multiples of \( k' \) such that the transition momenta are integer multiples of \( k' \) as expressed by Eq. (11). However, in the finite open Hubbard chains, for which no general statements on the functional form of the transition momenta can be made (see below), the character of the excitations can depend on \( U \). We suggest, therefore, that the transition momenta are given by

\[ k_q^c(U, N) = q\gamma_q(U)k_c(N) \]  \hspace{1cm} (13)

with \( k_c \) defined by Eq. (10), \( \beta \in \{ +, 1, -1, 4, -4 \} \), \( q = 1, 2, \ldots \), and \( \gamma_q(U) \geq 1 \). Values \( \gamma_q(U) > 1 \) can be interpreted as an effective shortening of the chains due to electron correlation.

Up to now the transition momentum quantum numbers \( q \) have been determined only for two low-lying ionic excitations in finite correlated Hubbard rings (see below and Ref. 53). The quantum numbers \( q \) associated with the transitions to the other ionic states and to the covalent states are still unknown because the Lieb-Wu equations have not yet been solved for these cases. For the linear chains no analytical results on quantum numbers and transition momenta are available since equations that are analogous to the Lieb-Wu equations for the rings have not been derived yet. In particular, there exists no rigorous justification for the suggestion in Eq. (13) concerning the transition momenta \( k_q^c \), such that this suggestion merely represents an attempt of a simple generalization of the \( U = 0 \) case. But for the moment, to avoid an unnecessary complication of the following discussion, we will assume that the finite chain transition momenta are independent of \( U \) and, hence, given by Eq. (12) instead by Eq. (13). Only later, for those cases, for which the generalization (13) cannot be avoided, we will use this generalization and try to determine, thereby, the scaling factor \( \gamma_q(U) \).

For \( U \neq 0 \) we would like to conjecture now that the dispersion relations when combined with the transition momenta provide approximate excitation energies for finite systems. Thus the excitation energies for finite Hubbard ring molecules are approximated by the composite functions \( E_{q'}^c = E_{q'}^c k_q' \) defined by

\[ E_{q'}^c(N) = E_{q'}^c \left( \frac{2\pi}{N} \right) \]  \hspace{1cm} (14)

and for open chains by the composite functions \( E_{q'}^c = E_{q'}^c k_q' \) defined by

\[ E_{q'}^c(N) = E_{q'}^c \left( \frac{\pi}{N + 1} \right) \]  \hspace{1cm} (15)

with \( \beta \in \{ +, 1, -1, 4, -4 \} \) and \( q = 1, 2, \ldots \). In case that the generalization (13) should apply to the chain transition momenta, the excitation energies should be approximately given by inserting the transition momenta \( k_q^c(U, N) \) instead of the momenta \( k_q^c(N) \) into the dispersion relations \( E^\beta \). Independently of the choice of the functional form of the chain momenta, however, those intervals of the dispersion relations \( E^\beta(k) \) that are linear in \( k \), should give rise to a linear behavior of the excitation energies for finite rings on a \( 1/N \) scale and for finite chains on a \( 1/(N + 1) \) scale. These conjectures will be investigated now.

\[ \text{B. Hubbard rings} \]

The excitation energies for the ionic states \( 1^1E_{1s}^+ \) and \( 1^1E_{2s}^+ \) of finite Hubbard rings with \( N = 4n + 2 \) atoms (see Sec. III for the nomenclature) have been determined by Hashimoto by an exact numerical solution of the Lieb-Wu equations. The excitation energies of the \( 1^1E_{1s}^+ \) state evaluated by Hashimoto for weak \( U/|t| = 1.25 \), medium \( (U/|t| = 2.08) \), and strong \( (U/|t| = 5) \) electron-electron repulsion are presented in Fig. 2 on a \( 1/N \) scale. It should be noted here that realistic polynern correspond to a \( U/|t| \) ratio of 1.5.

The transition momentum associated with the \( 1^1E_{1s}^+ \) excitation is \( 2\pi/N \), i.e., the momentum quantum number has the value \( q = 1 \). In order to test our suggestion that the dispersion relation \( E^+ \) of the conduction band of the infinite Hubbard chain provides an estimate of the low-lying ionic excitations of finite ring molecules we provide also in Fig. 2 the excitation energies \( E_1^+(N) \) resulting from Eq. (14).

Figure 2 demonstrates that the exact excitation energies of the \( 1^1E_{1s}^+ \) state for large \( N \) converge to the function \( E_1^+ \). In fact, in the cases of weak and intermediate electron-electron repulsion the exact excitation energies for all ring sizes follow closely the excitation energy function \( E_1^+ \) derived from the dispersion relation \( E^+ \). Be-
cause $E_1^+ \sim N^{-2}$ in the limit of large $N$, this agreement implies that also the excitation energies for large $N$ converge as $N^{-2}$ to the infinite chain optical gap. In the case of weak and intermediate electron-electron repulsion the size dependence actually behaves as $N^{-1}$ for rings with up to 100 atoms and assumes the $N^{-2}$ dependence only for larger rings. The latter deviation from the $N^{-1}$ dependence for weak and intermediate electron-electron repulsion is very small. Consequently, for an estimate $\Delta E_0$ of the optical gap in polycetylene with a known $U/|t|$ ratio of about 1.5, one can extrapolate the excitation energies of finite rings as $N^{-1}$, i.e., describe the excitation energies by the simple functional dependence

$$\Delta E = \Delta E_0 + \alpha/N$$

without introducing a large error. However, because of the asymptotic $N^{-2}$ dependence the gap energy $\Delta E_0$ in this equation will always underestimate the true gap energy $E^+(0)$.

The results in Fig. 2 represent only the lowest lying optically allowed $1^1E_{1u}^+$ state of Hubbard rings. In Fig. 3 we compare for intermediate electron-electron repulsion ($U/|t| = 2.08$) the excitation energies of the $1^1E_{1u}^+$ state with those of the $1^1E_{2g}^0$ state. The $1^1E_{2g}^0$ state is the third ionic state in Hubbard rings and according to Hashimoto $^5$ is characterized by a value $q = 2$ of the transition momentum quantum number. (The second ionic state, which is not shown in Fig. 3, is of $1B_{1u}^+$ symmetry. Due to the degeneracy of the orbital pattern of the $N = 4n + 2$ rings this state is nearly degenerate with the $1^1E_{1u}^+$ state. $^4$ Unfortunately, no exact results on the $1^1B_{1u}^+$ state are available up to now). The size dependence of the exact excitation energies has been fitted for both ionic states to expressions of the type (16). The energy gaps $\Delta E_0$ resulting for the two transitions are identical (0.29 eV) and underestimate the exact value of $E^+(0) = 0.48$ eV. The slopes $\alpha$ for the two excitations differ by a factor of 1.95. This value is close to the value $q = 2$ expected from predictions based on the dispersion relation $E^+$. These predictions assign the excitation energies $E_1^+(N)$ to the $1^1E_{1u}^+$ state and $E_2^+(N)$ to the $1^1E_{2g}^0$ state. The close to linear $1/N$ behavior of $E_1^+$ and $E_2^+$ in the range $10 < N < 50$ implies that the slopes $\alpha$ in Eq. (16) yield information on the momentum quantum numbers $q$ to be attributed to the electronic excitations. Thus, the energies for ionic excitations of Hubbard rings actually follow the approximate law

$$\Delta E = \Delta E_0 + q\alpha/N, \quad q = 1, 2, \ldots$$

We will consider the size dependence of the excitation energies involving the covalent states. Figure 4 provides the excitation energies of the two lowest covalent singlet states $1^1B_{1u}^0$ and $1^1E_{2g}^0$ of finite Hubbard rings in case of an intermediate electron-electron repulsion described by a $U/|t| = 2$ ratio. For comparison Fig. 4 also shows the excitation energies of the lowest ionic singlet state $1^1E_{1u}^0$. In lieu of an analytical description these excitation energies had to be evaluated by our MRD-CI method, for
which reason only data points for rings with 6, 10, and 14 atoms are available. Such limited information does not yet allow a convincing demonstration of the asymptotic behavior of coherent ring excitations. Nevertheless, the following analysis of the data shown in Fig. 4 seems to indicate that the coherent excitations follow the same pattern as the ionic excitations.

The analysis starts from a comparison of the MRD-CI excitation energies of the $1^1B_{1u}$ and $1^1E_{2g}$ states with predictions of these energies derived from the dispersion relations $E_j^{\pm}$. For small $k$ all dispersion relations $E_j^{\pm}(k)$, $j = 1, 2, 4$, of coherent excitations eventually become linear and the linear range extends towards larger momenta $k$ for larger values of $j$. Thus, all ring excitation energy functions $E_j^{\pm}$ become linear for large $N$ on a $1/N$ scale and the linear range extends further towards small ring sizes $N$ for small momentum quantum numbers $q$ and large values of $j$. As shown in Fig. 4 the linear range of the ring excitation energy function $E_j^{\pm q}$ ($q = 1, j = 1, 3$) extends up to about $N = 10$. The excitation energies of the lowest coherent state $1^1B_{1u}$ for large $N$ approach this function from above. The tangent of the function $E_j^{\pm}$ at $1/N \to 0$, that, according to the arguments given above, represents an even closer approximation to the function $E_j^{\pm}$ (not shown in Fig. 4), provides an excellent approximation to the $1^1B_{1u}$ excitation energies for rings of all the sizes considered. Multiplication of the slope of the tangent by a factor 2, which according to Eqs. (11) and (14) yields the tangent to the function $E_j^{\pm q}$ ($q = 2$), provides an asymptotic limit of the excitation energies of the second coherent state, the $1^1E_{2g}$ state. We expect, therefore, that the coherent excitations follow also the simple law (17) with $q = 1$ for the $1^1B_{1u}$ state and $q = 2$ for the $1^1E_{2g}$ state. This important observation, so far only vaguely indicated by the results in Fig. 4, will be demonstrated more convincingly for open Hubbard chains and for polyenes described by the PPP model.

The close agreement between the tangent of $E_j^{\pm}$ at $1/N \to 0$ and the MRD-CI excitation energies of the lowest coherent state suggests that the extrapolation of the MRD-CI excitation energies on a $1/N$ scale for rings should be a reliable tool for the determination of the coherent gap $E^{\pm}(0)$. If sufficient data on large systems and on higher excited states are available such extrapolations according to (17) can also yield the coherent momentum quantum numbers $q$.

C. Linear Hubbard chains

We want to demonstrate now that the ionic and coherent excitations in linear Hubbard chains exhibit a size dependence of their excitation energies which is very similar to that of the ring compounds as described by Eq. (17). There exist, however, two important differences. (1) Since the chain momenta $k_j^c$ exhibit a $1/(N + 1)$ behavior instead of the $1/N$ behavior of the ring momenta, the size dependence of the chain excitation energies has to be considered on the appropriate $1/(N + 1)$ scale. (2) The transition momenta $k_j^c$ of the ionic excitations turn out to be noninteger multiples of the elementary chain momentum $k$ with a scaling factor $\gamma$ depending on the degree of electron-electron repulsion.

First we want to show that Eq. (13) instead of Eq. (12) applies to the transition momenta $k_j^c$ and that the factor $\gamma$ in Eq. (13) assumes values $\gamma_j^c(U) > 1$ if $U/|t| > 1$. For this purpose we compare in Fig. 5 the excitation energies of the lowest $(q = 1)$ ionic states in rings $(1^1E_{2u})$ and linear chains $(1^1B_{1u})$ with the chain excitation energy functions $E_j^{\pm}$, $q = 1, 2$. These functions have been derived from the dispersion relations $E^{\pm}$ according to Eq. (15) which does not take a value $\gamma_j^{c+}(U) > 1$ for $U/|t| > 0$ into account.

In the case of vanishing Coulomb repulsion $(U/|t| = 0)$ the analytical solutions for the $1^1E_{2u}$ and $1^1B_{1u}$ excitation energies are given by $E_j^{c+}(N)$ and $E_j^{c-}(N)$, respectively. According to Eqs. (8)–(10), (14), and (15) these functions are

$$E_j^{c+}(N) = 4t \sin(\pi/N)$$

for the ring $(1^1E_{2u})$ states and

$$E_j^{c+}(N) = 4t \sin(\pi/2(N + 1))$$

for the linear chain $(1^1B_{1u})$ states. Since the ring momentum $k_j^c$, given by $(9)$, is about two times larger than the chain momentum $k_j^c$, the ring excitation energies increase about two times faster than the chain excitation energies with decreasing molecular size $N$. This is exhibited in Fig. 5 by the fact, that the ring $1^1E_{2u}$ excitation energies
FIG. 5. The end effect on the excitation energies of the lowest ionic singlet state of the Hubbard model and its dependence on electron correlation. The dispersion relations $E^+$ for the particle-hole singlet excitation of the infinite chain evaluated at one ($E^+_1$) and two ($E^+_2$) quanta of the finite chain momentum $k^s=\pi/(N+1)$ [cf. Eq. (15)] are compared with the exact energies of the $E^+_1$ state (squares) of the finite open chains and of the $E^+_2$ state (double crosses) of the rings in the case of the uncorrelated electron system ($U/|t|=0$) and with MRD-CI excitation energies of the $E^+_2$ state in the case of a strongly correlated electron system.

$q=1$, given by Eq. (18), are located close to the excitation energies $E^+_2$ which are associated with the transition momenta $2k^s$.

The difference between the energies $E^+_1(N)$ and $E^+_1(N+1)$ has to be attributed to the "end effect." Equations (18) and (19) demonstrate that the end effect is large. This result is in variance with Hashimoto's treatment of the end effect$^{53}$ that has been based on first-order perturbation theory. This author determined that the end effect vanishes faster than $1/N$ which is obviously not true in the $U=0$ limit [cf. Eqs. (8)--(10)].

However, the end effect becomes smaller when the degree of electron-electron interaction increases. This can be concluded from a comparison in Fig. 5 of the excitation energy of the $E^+_1$ state ($q=1$) with the excitation energy functions $E^+_1$ and $E^+_2$ for $U/|t|=4$. The former function would provide an estimate for the $E^+_1$ excitation energies if the transition momentum $k^s$ would be solely determined by the geometry, i.e., given by Eq. (10). That is not the case. However, the latter function which is close to the ring excitation energies $E^+_1(N)$ lies also close to the $E^+_2$ excitation energies. This implies that the chain momentum $k^s$ associated with the transition to the $E^+_2$ state in the case $U/|t|=4$ is about two times larger than the momentum $k^s=k^s$ associated with this transition in the case $U/|t|=0$. Consequently, Eq. (13) instead of Eq. (12) has to be employed for the description of the transition momenta of the ionic chain states and the factor $\gamma^+(U)$ increases from a value $\gamma^+(0)=1$ to the value $\gamma^+(4|t|)=2$.}

FIG. 6. The end effect on the excitation energies of the lowest covalent singlet state of the Hubbard model and its dependence on electron correlation. For the cases of vanishing ($U/|t|=0$) and of strong ($U/|t|=4$) electron correlation the lower edge $E^1$ of the dispersion relations of the singlet spin excitations in the infinite chain as well as the upper edge $E^4$ of these relations for excitations characterized by four holes in the $\lambda$ distribution (Ref. 37) have been evaluated at two quanta of the finite chain momentum $k^s=\pi/(N+1)$ resulting in the functions $E^1$ and $E^4$ [cf. Eq. (15)]. In the case of strong electron correlation these functions are compared with the MRD-CI excitation energies of the $A^-$ state of open chains, whereas in the case of vanishing electron correlation they are compared with the exact excitation energies of the $B_{1u}$ state of the rings (double crosses) and of the $A_g$ states of the open chains. The latter states are characterized by a double excitation from the highest occupied to the lowest unoccupied molecular orbital (large crosses) and by a single excitation from the highest occupied to the second to lowest unoccupied orbital (small crosses), respectively.

Figure 6 compares again for the cases $U/|t|=4$ and $U/|t|=0$ the excitation energies of the lowest covalent states $1B_{1u}$ in the ring ($q=1$) and $2A_g$ in the linear chain ($q=2$) with excitation energy functions derived according to Eq. (15) from the dispersion relations $E^1$ and $E^4$ assuming $q=2$. Here, the value $q=2$ for the $2A_g$ state derives from the fact that this state involves an excitation which promotes either a single electron across two single electron level spacings or promotes two electrons from the highest occupied to the lowest unoccupied single electron level, i.e., promotes two electrons across a single level spacing. As one can show for the case $U=0$, the crossing of each electron level spacing adds a minimum momentum value $k^s$ to the many electron transition.

For the case $U=0$ the ring and chain excitation energies are very close and asymptotically approach the $E^1$ and the $E^4$ excitation energy functions. The latter function represents the exact solution for the two-electron excitation (large crosses in Fig. 6) in linear chains. The two times larger quantum number of the lowest covalent excitation in the chains as compared to that in the rings.
and the little more than two times smaller value of the elementary chain momentum $k^c$ as compared to that of the ring momentum $k^r$ explain why a large, 1/$N$-type end effect is missing in the case of spin-wave excitations.

The excitation energies of the lowest covalent state for the $U/|t| = 4$ case are seen in Fig. 6 to lie below the corresponding excitation energies for the $U/|t| = 0$ case, i.e., they exhibit an ordering which is opposite to that of the ionic states for the same difference in the $U/|t|$ ratios (cf. Fig. 5). This demonstrates the well-known fact that electron correlation effects decrease the excitation energies of ionic states and increase the excitation energies of ionic states. The excitation energies, which were determined by means of the MRD-CI method, asymptotically approach the $E_1^{1-}$ and $E_2^{4-}$ excitation energy functions. This behavior indicates that the momentum changes associated with the ionic states in the linear chain compounds are not affected by the degree of electron-electron repulsion, i.e., that $\gamma^{-}(U) \equiv 1$ and that Eq. (12) applies to these transition momenta.

We want to consider now in further detail the size dependence of the covalent and the ionic excitations in linear Hubbard chains. For this purpose we present in Figs. 7 and 8 the excitation energies for the case $U/|t| = 2$ involving the two lowest covalent and ionic excitations.

![FIG. 7. Linear 1/(N+1) extrapolations of the Hubbard-MRD-CI excitation energies of the two lowest covalent singlet excitations of finite linear polyenes and comparison with corresponding excitation energies in the infinite chain for the case of intermediate electron correlation characterized by the value $U/|t| = 2$ of the Hubbard parameter. The extrapolation (lower dashed line) of the $2^{1}A_{u}^{-}$ excitation energies (crosses) represents a least square deviation fit according to Eq. (20) to the MRD-CI results obtained for the $N = 10, 12$, and 14 $\pi$-electron polyenes, whereas the extrapolation (upper dashed line) for the $1^{1}B_{u}^{+}$ state (double crosses) is the line connecting the excitation energies at $N = 12$ and 14. The slope of the fit to the $2^{1}A_{u}^{-}$ excitation energies ($q = 2$) is $\alpha = 12.15$ eV and the ratio of the slopes of the two extrapolations is 1.46. For comparison with the $2^{1}A_{u}^{-}$ excitation energies the dispersion relations $E_{1}^{1-}$ and $E_{2}^{4-}$ of the infinite system have been evaluated at two quanta of the finite chain momentum $k^c = \pi/(N + 1)$ (see text and Fig. 5 caption for further explanation).](image)

![FIG. 8. Linear 1/(N+1) extrapolations of the Hubbard-MRD-CI excitation energies of the two lowest ionic singlet excitations of finite linear polyenes and comparison with corresponding excitation energies in the infinite chain for the case of intermediate electron correlation characterized by the value $U/|t| = 2$ of the Hubbard parameter. As a reference the excitation energies (crosses) and extrapolation (dashed line) for the ionic $2^{1}A_{u}^{-}$ state are also shown. The extrapolation (lower solid line) of the $1^{1}B_{u}^{+}$ excitation energies (circles) represents a fit according to Eq. (20) to the MRD-CI results obtained for the $N = 8, 10, 12$, and 14 $\pi$-electron polyenes, whereas the extrapolation (upper solid line) for the $1^{1}A_{u}^{+}$ state (squares) is the line connecting the excitation energies at $N = 12$ and 14. The ratio of the slopes of the two extrapolations is 1.67, the extrapolated optical gap is $\Delta E_{\text{opt}} = 0.29$ eV and the slope of the fit to the $1^{1}B_{u}^{+}$ excitation energies ($q = 1$) is $\alpha = 24.11$ eV. For an evaluation of the end effect the $1^{1}B_{u}^{+}$ and $1^{1}A_{u}^{+}$ excitation energies are compared with corresponding excitation energies of the infinite system calculated from the dispersion relation $E_{1}^{+}$ at one ($E_{1}^{+}(1)$) and two ($E_{1}^{+}(2)$) quanta of the finite chain momentum $k^c = \pi/(N + 1)$.](image)
2 \textsuperscript{1} A_{\text{g}^-} \text{ state} the latter value corresponds to a momentum quantum number \( q = 3 \) for the 1\textsuperscript{1} B_{\text{u}^-} \text{ excitation}. A generalization of this finding suggests that there is a class of covalent excitations in linear Hubbard chains which for large \( N \) approximately follows the law

\[
\Delta E = \Delta E_0 + q \alpha / (N + 1) \tag{20}
\]

with \( q = 2, 3, 4, \ldots \). Equation (20) is analogous to the law for rings expressed by Eq. (17). The close to linear 1/\( N + 1 \) behavior of the chain excitation energies will actually be documented in further detail for the PPP model in the next section.

The results in Fig. 7 on the covalent states show that one can extrapolate indeed numerical excitation energies for finite chains to the infinite chain limit by means of a 1/\( N + 1 \) size dependence if these extrapolations are based on excitation energies of compounds comprising 10 sites and more.

Figure 8 presents the size dependence of the excitation energies involving the two lowest lying ionic states. The excitation energies of the 2 \textsuperscript{1} A_{\text{g}^+} \text{ state} are also included in Fig. 8 to demonstrate for the U/|t| = 2 linear Hubbard chain that this state lies energetically below the optically allowed ionic states.

The excitation energies of the ionic state 1\textsuperscript{1} B_{\text{u}^+} \text{ (q = 1)} are compared in Fig. 8 with the excitation energy functions \( E_{\text{g}^+}(q) \) and \( E_{\text{b}^+}(q) \) that correspond to momentum quantum numbers \( q = 1 \) and \( q = 2 \) according to Eq. (15). The results show that for the U/|t| = 2 case the 1\textsuperscript{1} B_{\text{u}^+} \text{ excitation carries} a momentum \( k_{\text{g}^+} \approx 1.5k_s \), i.e., \( \gamma_{\text{g}^+}(2|t|) \approx 1.5 \) a value which is halfway between that for the U=0 case and the U=4|t| case.

It is of interest to determine the transition momentum \( k_{\text{g}^+} \) associated with the second ionic excitation involving the 1\textsuperscript{1} A_{\text{g}^+} \text{ state. The ratio of the slopes of the linear 1/\( N + 1 \) extrapolations shown in Fig. 8 for the two ionic states is 1.67. If \( k_{\text{g}^+} \) would be given according to a relation like \( k_q = qk_{\text{b}^+}, \text{i.e., if the factor} \gamma_q(U) \text{ in Eq. (13)} \) would be independent of \( q \), one would expect a ratio of 2. There are two possible explanations for this difference. (1) The difference may be due to an accidental addition of the errors inherent to our MRD-CI description\textsuperscript{13} and to our linear extrapolations. Thus, it may be that a ratio closer to the value 2 would result if the extrapolations were based on exact excitation energies for longer compounds. (2) The deviation of the ratio 1.67 from the value 2 indicates a true physical effect, i.e., \( \gamma_q(U) \text{ decreases with increasing} q \). The value 1.67 implies then \( \gamma_q(2|t|) \approx 1.25 \). The U dependence of the transition momentum \( k_{\text{g}^+} \) as well as its decrease with increasing \( q \) could be due to the development of an excitonic character of the lowest energy ionic excitations in a linear chain geometry. Such character would imply that the particle-hole binding energy decreases for decreasing electron correlation. Furthermore it could also imply that higher excitations are less affected by this excitonic binding energy, i.e., that \( \gamma_q(U) \rightarrow 1 \text{ for both} U \rightarrow 0 \text{ and} q \rightarrow \infty \). Then also for the ionic transitions Eq. (12) instead of Eq. (13) should hold if higher excitations are involved and if electron correlation is small.

Figure 8 demonstrates that the linear dependence in 1/\( N + 1 \) is satisfied very well by MRD-CI excitation energies for the ionic states. This suggests that finite size energies can be extrapolated to the infinite chain limit for ionic states as well. However, the resulting value \( \Delta E_0 = 0.29 \text{ eV} \) for the optical gap is too small compared to the exact value \( E^+ (0) = 0.41 \text{ eV} \). The error in the gap energy should be less for calculations of realistic polyynes since these systems correspond to a U/|t| = 1.5 ratio for which the \( N + 1 \)\textsuperscript{-2} deviation from the 1/\( N + 1 \) extrapolation is less significant.

V. POLYENE AND POLYACETYLENE ELECTRONIC SPECTRA IN THE PPP MODEL

We want to show in this section that close similarities exist between linear polyynes as described by the Hubbard and by the PPP Hamiltonian. The similarities are found for the asymptotic size dependence of the ionic and covalent transitions. We will further investigate in this section the effect of bond alternation on the spectra of polyenes, in particular, in the limit of the infinite polyene, i.e., polyacetylene (CH)n. Lastly, we consider to what extent dielectric screening of the electron-electron repulsion influences the spectra of polyenes and polyacetylene.

In order to apply the results of Sec. IV we compare the strength of the electron-electron repulsion in the PPP and in the Hubbard model. For this purpose one has to determine the ratio \( (U_{ij} - U_{ij+1})/|t| \), where \( U_{ij} \) is the Coulomb integral for electrons at sites \( i \) and \( j \), and \( t \) is the resonance integral given by Eq. (5). Our parametrization of the PPP Hamiltonian yields the value 1.53 for this ratio. We conclude that the short-range Coulomb repulsion between the \( \pi \) electrons is weaker in the PPP model than in a Hubbard model with a U/|t| = 2 ratio.

A second measure for the effect of the Coulomb repulsion in the PPP and the Hubbard model is the ground state correlation energy per site. This energy is defined as the energy difference between the SCF ground state and the ground state as described by the MRD-CI method divided by the number of carbon atom. The ground state correlation energy per site measures about 0.15 eV for the PPP model with bond alternation, 0.18 eV for the PPP model without bond alternation,\textsuperscript{13} and 0.25 eV for the U/|t| = 2 Hubbard model. This comparison indicates, too, that correlation effects are weaker in the PPP model than in the U/|t| = 2 Hubbard model. Consequently, the results on the Hubbard model let us expect that the size dependence of the spectra of the polyenes in the PPP description should obey a 1/\( N + 1 \) dependence as expressed by Eq. (20) for compounds with 10 carbons and more. Such dependence, actually a 1/N dependence, had been observed and predicted before for the optical transition to the ionic 1\textsuperscript{1} B_{\text{u}^+} \text{ state} of polyynes.\textsuperscript{30,35}

A. Nonalternating model polyenes

In the Hubbard model of linear polyenes as described by the Hamiltonian (1) all resonance integrals \( t \) are chosen to be identical. This implies according to Eq. (5) that all bond lengths are assumed to be equal. We will, therefore,
consider the PPP model of polyenes without bond alternation first, since that should yield the closest similarities to the results in the previous section.

Figure 9 presents our MRD-CI excitation energies for the two lowest ionic and covalent excitations in nonalternating polyenes. The calculations were carried out for polyenes with \( N = 4, 6, \ldots, 14 \) carbon atoms. The results in Fig. 9 show that the excitation energies for \( N = 10, 12, 14 \) follow closely a \( 1/(N+1) \) dependence. The gap energies and slopes resulting from a fit according to Eq. (20) to the lowest ionic \((q = 1)\) and covalent \((q = 2)\) transitions are provided in Table I. In agreement with the expectation voiced in the paragraph above, the \( 1/(N+1) \) size dependence holds well for polyenes with 10 and more carbons.

It is a very interesting result of our study that the energy gap for the covalent transitions vanishes in the limit \( N \to \infty \) for the PPP polyene without bond alternation, i.e., this well-known feature of the Hubbard model is not altered when the range of the Coulomb repulsion is extended in the PPP model. A comparison of Fig. 9 and of the data in Table I with the results for the \( U/Lt = 2 \) Hubbard model in Figs. 7 and 8 shows also that the slopes of the extrapolated size dependences of the covalent transitions are similar for the two models. The similarity between the ionic transitions is not as striking. The asymptotic optical gap is much larger in the PPP model than in the \( U/Lt = 2 \) Hubbard model proving that also long-range electron interactions contribute to this gap. However, the slopes of the respective extrapolations and, in particular, the ratios of these slopes are similar such that the analysis of these ratios, that has been carried out in detail for the \( U/Lt = 2 \) Hubbard model and has lead to the determination of transition momenta and quantum numbers, applies equally well to the nonalternating PPP model considered here.

B. Spectra of alternating polyenes

We now turn to the most realistic model of polyenes and polyacetylene that is based on the PPP Hamiltonian and on a geometry with bond alternation, the latter being described by \( \delta = 0.10 \) Å (see Sec. III). The results of the corresponding MRD-CI calculations are the most relevant in this paper. For this reason, the calculations were carried out to the largest degree of completion, i.e., they cover polyenes with 4 to 16 carbons and include three covalent \((2^1 A_1^+, 1^1 B_1^-, 3^1 A_2^-)\) and two ionic \((1^1 B_2^+, 1^1 A_2^+)\) states. The size dependence of the respective excitation energies is presented in Fig. 10.

The overall appearance of the spectra in Fig. 10 is similar to that of linear Hubbard chains. There are two bands of excitations, the covalent band and the ionic band. The size dependence of the excitation energies in both bands follows an \( 1/(N+1) \) behavior as illustrated in Fig. 10. We have exhibited the \( 1/(N+1) \) dependence by fitting straight lines through the data points in Fig. 10. The lines have been obtained as follows: We have first drawn the two lines through the data points for the primary, i.e., lowest, covalent state and the primary ionic state for polyenes with 12, 14, and 16 carbons. These lines correspond to a size dependence as described by the function in Eq. (20) with \( q = 2 \) for the \( 2^1 A_1^+ \) state and \( q = 1 \) for the \( 1^1 B_2^+ \) state. The corresponding slopes \( \alpha \) and the asymptotic gaps \( \Delta E_0 \) are presented in Table I. We have then replotted the function describing the primary covalent excitation for \( q \) values 3 and 4. The resulting lines are found in Fig. 10 to coincide asymptotically with the excitation energies of the \( 1^1 B_2^- \) and of the \( 3^1 A_2^- \) state, respectively.

Since the analysis for the Hubbard rings and chains revealed that the \( 2^1 A_1^+ \) excitation carries the momentum \( 2k^c \), \( k^c \) defined in (10), this latter observation implies that the \( 1^1 B_2^- \) excitation carries a momentum \( 3k^c \) and the \( 3^1 A_2^- \) excitation carries a momentum \( 4k^c \), proving thereby our conjectures in Sec. IV. We have also replotted the function describing the primary ionic excitation—up to a very small shift of the asymptotic gap—with a \( q \) value of

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</table>

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**TABLE I.** Parameters of extrapolations. The parameters of the \( 1/(N+1) \) extrapolations [see Eq. (20)] have been obtained for the various PPP models of polyenes from fits to MRD-CI excitation energies of the primary covalent and ionic states as shown in Figs. 9, 10, and 12.
The resulting line falls asymptotically on the excitation energies for the \(1^1A_u^+\) state implying that this excitation carries a momentum two times larger than that of the \(1^1B_u^+\) state. The value 2 determined for the ratio of the slopes of the ionic-state extrapolations shown in Fig. 10 is a little surprising in view of the analysis of the momenta associated with the ionic transitions in the Hubbard model. This analysis had revealed that these momenta increase with increasing interelectronic forces and, possibly, decrease with increasing quantum numbers \(q\). The latter decrease had furnished a possible explanation of the rather small ratio 1.67 determined for the ionic-state extrapolations in the \(U/|t|=2\) Hubbard model and in the PPP model without bond alternation. The value 2 found here in the PPP model of alternating polyenes may imply either that alternation decreases the elementary chain momentum of ionic transitions from the value \(k_{\pm}^\pm \approx 1.5k^e\) to the value \(k_{\pm}^\pm \approx k^e\), i.e., that \(\gamma_{\pm}(U)\approx 1\) [cf. Eq. (13)], or else that there is no strong dependence of \(\gamma_{\pm}(U)\) on \(q\) in alternating polyenes.

One of the most interesting results shown in Fig. 10 is the prediction of a finite asymptotic gap of the covalent states which measures \(E^{-}(0)=1.85\) eV. This gap has to be attributed solely to bond alternation. With the value for \(\alpha\) given in Table I one obtains from Eqs. (10), (12), (15), and (20) a linear approximation to the dispersion relation of the covalent excitations in the infinite alternating polyene that is given by

\[
E^-(k) = E^-(0) + \alpha k
\]

with \(\alpha = \alpha/\pi = 2.75\) eV. The gap of ionic excitations which corresponds to the gap of the conduction band in \((\text{CH})_x\) is found to be increased by a third over that of the polyene without bond alternation (see Fig. 9). Since single electron theories predict a vanishing optical gap for nonalternating polyenes, this implies that, contrary to the behavior of the covalent gap, only about 30% of the optical gap in \((\text{CH})_x\) should be due to Peierls distortion, the major fraction being due to electron correlation.\(^12\)

C. The origin of the covalent gap and the spectrum of triplet excitations

Before we attempt to relate the results shown in Fig. 10 and Table I to observed properties of \((\text{CH})_x\) (see Sec. V D) we would like to give a physical interpretation of the covalent gap \(E^{-}(0)\) predicted for \((\text{CH})_x\). For this purpose we first note that the alternation of short and long C-C distances observed for the ground state of \((\text{CH})_x\) (Ref. 56) indicates the existence of stable \(\pi\) bonds. Thus, the ground state of \((\text{CH})_x\) can be represented as a sequence of ethylene units each characterized by a singlet spin pairing of the \(\pi\) electrons (see Fig. 11(a)).

The \(2^1A_g^-\) state has been characterized as involving two intraethylene triplet excitations which are coupled to an overall single state.\(^13,14\) In fact, the excitation energy of the primary covalent excitation measures about twice the excitation energy of the lowest triplet excitation. A typical spin coupling diagram contributing to the \(2^1A_g^-\) state is shown in Fig. 11(b). Two of the singlet pairings of the ground-state structure in Fig. 11(a) have been converted to a triplet pairing. The location of the triplet pairing

\[
\text{(a) } 1^1\tilde{A}^-
\]

\[
\text{(b) } 2^1\tilde{A}^-
\]

 FIG. 11. Schematic representation of the electronic structure of the ground state (a) and of the first excited covalent state (b) of the polyenes. (a) The ground-state structure of alternating single and double bonds shown in \(\alpha\) is represented by the singlet spin pairings within the ethylene units depicted in \(\beta\). (b) As indicated in \(\beta\) the excited state is created by simultaneous spin flips in two ethylene units, the resulting triplet pair being coupled to an overall singlet. Each intraethylene triplet corresponds to a single rather than to a double bond as sketched in \(\alpha\).
is not fixed but one has to imagine that the triplet excitations are moving over the whole polyene.

We would like to demonstrate now that the above characterization of the $2^1 A_g$ state as a triplet-triplet ($TT$) excitation can be generalized to the whole class of low-lying covalent singlet excitations. This conjecture implies that one should obtain the excitation energies of the higher covalent singlet states ($1^1 B_u$, $3^1 A_g$, etc.) by adding the excitation energies of those covalent triplet excitations which couple to form the respective $TT$ state.

A polyene of $N$ carbon atoms can accommodate as many linearly independent covalent triplet states as there are ethylene units, i.e., $N/2$. For the $N=10$ polyene the PPP excitation energies $\Delta E_{\text{FCI}}$ of the three lowest covalent triplet states ($1^1 B_u$, $1^1 A_g$, $2^1 B_u$) are given in Table II. These energies have been calculated by Soos and Ramasesha using a complete valence-bond description. By forming irreducible products of the triplets one obtains the symmetry labels of the $TT$-excitations, e.g., $1^1(3 B_u \otimes 1 B_u)^{-1} A_g$, and by adding the corresponding energies one obtains an estimate $\Delta E_{\text{sum}}$ for the excitation energies of the $TT$ states. Table III gives for the five lowest covalent singlet excitations in the $N=10$ polyene, the decomposition into triplet excitations, and compares the estimates $\Delta E_{\text{sum}}$ with exact ($\Delta E_{\text{FCI}}$) and close to exact ($\Delta E_{\text{OCl}}$) computational results. Again, the exact excitation energies have been taken from Ref. 12 whereas the close to exact excitation energies have been calculated by our Q-CI method.

As exhibited by Table III the sums $\Delta E_{\text{sum}}$ of the triplet excitation energies furnish a close approximation to the excitation energies of the $TT$ states indicating that our generalization is actually correct. The approximation is a little worse for the $3^1 A_g$ and $4^1 A_g$ states which should be (nearly) degenerate according to the estimate $\Delta E_{\text{sum}}$. These states show a splitting of $\approx 0.4$ eV centered around the $\Delta E_{\text{sum}}$ value. For a finite compound like the $N=10$ polyene considered here such splitting had to be expected because the two states have the same symmetry. The $N$ dependence of the excitation energies shown in Ref. 13 leads to the expectation that, in fact, the $3^1 A_g$ and $4^1 A_g$ states are degenerate in the infinite system. The fifth covalent singlet excitation $2^1 B_u$ is characterized in Table III as the irreducible product $1^1(3 B_u \otimes 1^3 B_u \otimes 1^3 B_u)$ of the lowest triplet state and, consequently, should have three times the excitation energy of this state. Hence, the $2^1 B_u$ state is suggested to belong to a different class of covalent singlet excitations—the class of $TTT$ states—which, as we will show below, should extrapolate to a covalent gap different from that of the $TT$ states. The characterization of the $2^1 B_u$ state as a $TTT$ state as well as that of the other covalent singlets as $TT$ states has been checked by us also for the $N=12$ polyene by means of comparisons similar to those in Tables III.

The characterization of the low-lying covalent singlet excitations as $TT$ states suggests that one should obtain the dispersion relation $E_T(k)$ of the covalent triplet states in the infinite alternating polyene from that of the $TT$ states, given in Eq. (21), according to

$$E_T(k) = E_{\text{FCI}} - (0)/2 + \alpha k. \tag{22}$$

Generalizing the relationship between infinite chain dispersion relations and finite chain excitation energies expressed by Eq. (15) to triplet excitations we conjecture that the dispersion relation $E_T(k)$ provides an approximation $E_T^N(k) = E_T(q_\Xi k^N)$, $q_\Xi = 1, 2, \ldots, N/2$, for the excitation energies of covalent triplet states. Table II demonstrates for $N=10$ that indeed the approximate excitation energies $E_T^N(10)$ compare very closely with the exact values $\Delta E_{\text{FCI}}$ for $q_\Xi = 1, 2, 3$.

An intraethylene triplet excitation involves the transformation of an ethylene double bond to a single bond [cf. Fig. 11(b)]. Qualitatively one can interpret the triplet gap

<table>
<thead>
<tr>
<th>Triplet</th>
<th>$\Delta E_{\text{FCI}}$</th>
<th>$E_T(10)$</th>
<th>$q_T$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$1^3 B_u$</td>
<td>1.76</td>
<td>1.71</td>
<td>1</td>
</tr>
<tr>
<td>$1^1 A_g$</td>
<td>2.57</td>
<td>2.49</td>
<td>2</td>
</tr>
<tr>
<td>$2^1 B_u$</td>
<td>3.39</td>
<td>3.28</td>
<td>3</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Singlet</th>
<th>$\Delta E_{\text{OCl}}$</th>
<th>$\Delta E_{\text{FCI}}$</th>
<th>Triplet-triplet</th>
<th>$\Delta E_{\text{sum}}$</th>
<th>$E_{qs}(10)$</th>
<th>$q_S = q_T + q_{\Xi}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$2^1 A_g$</td>
<td>3.42</td>
<td>(3.40)</td>
<td>$1^1 B_u \otimes 1^3 B_u$</td>
<td>3.51</td>
<td>3.42</td>
<td>$2 = 1 + 1$</td>
</tr>
<tr>
<td>$1^1 B_u$</td>
<td>4.23</td>
<td>(4.23)</td>
<td>$1^3 B_u \otimes 1^1 A_g$</td>
<td>4.32</td>
<td>4.20</td>
<td>$3 = 1 + 2$</td>
</tr>
<tr>
<td>$3^1 A_g$</td>
<td>4.93</td>
<td>(—)</td>
<td>$1^1 A_g \otimes 3 B_u$</td>
<td>5.13</td>
<td>4.98</td>
<td>$4 = 2 + 2$</td>
</tr>
<tr>
<td>$4^1 A_g$</td>
<td>5.31</td>
<td>(—)</td>
<td>$1^3 B_u \otimes 2 B_u$</td>
<td>5.14</td>
<td>4.98</td>
<td>$4 = 1 + 3$</td>
</tr>
<tr>
<td>$2^1 B_u$</td>
<td>5.34</td>
<td>(5.32)</td>
<td>(1 $1^1 B_u)$</td>
<td>5.27</td>
<td>5.13</td>
<td>$3 = 1 + 1 + 1$</td>
</tr>
</tbody>
</table>
$E^{-}(0)/2$ in Eq. (22) as the energy required to remove one of the distinct $\pi$ bonds which make up the ethylene units of an infinite alternating polyene. The ground state of a nonalternating polyene is degenerate. Thus there are no energetically favored singlet spin pairings and the triplet gap vanishes.

The slope $\alpha$ of the dispersion relation $E^{T}(k)$ furnishes the group velocity $v_T$ at which an intraethylene triplet excitation moves along the polyacetylene chain. $v_T$ is given by $\hbar \alpha /\hbar$, where $\hbar = 1.4\ \text{A}$ is the average C--C distance, and measures $0.58 \times 10^6 \text{ m/s}$. The data on the slope $\alpha$ of the $1/(N+1)$ extrapolations in Table I show that $v_T$ increases with decreasing bond alternation and from the discussion of the Hubbard model in Sec. IV one finds that $v_T$ decreases with increasing electron correlation.

In the analysis presented above the properties of the triplet states have been deduced from a decomposition of the covalent singlet excitations. Conversely, the results on the triplets can now be used to obtain new insights into the properties of the covalent singlets. The covalent gap appears then as the energy required to break the $\pi$ bonds in two of the ethylene units of (CH)$_2$. Furthermore, the quantum numbers $q_S = 2, 3, 4, \ldots$ associated with the $TT$ excitations are explained as the sums

$$q_S = q_T + \bar{q}_T$$  \hspace{1cm} (23)

of the quantum numbers $q_T$ and $\bar{q}_T$ of the associated triplet excitations. Analogously the quantum numbers of the $TTT$ excitations are given by $q_S = q_T + 2\bar{q}_T + \bar{q}_T$ and with Eqs. (22), (10), and (12) the $N$ dependence is given by the law

$$E^{TTT}_{qs} - (N) = 3E^{-(0)}/2 + aq_S/N(N+1).$$  \hspace{1cm} (24)

We have checked the correctness of Eq. (24) for the lowest $TTT$ excitation $2^1B_{uv}^-$ using Q-CI results for $N = 6, 7, \ldots, 12$. Table III shows for the $N = 10$ polyene that, indeed, the $TT$ state quantum numbers $q_T + \bar{q}_T$ obtained by combination are identical to the values $q_S$ obtained from the slopes of the extrapolations in Fig. 10 and that the estimates $E^{TTT}_{qs}$ obtained according to Eq. (20) for the $TT$ excitations and $E^{TTT}_{qs}$ obtained according to Eq. (24) for the $TTT$ state are very close to the exact energies.

The composition principle expressed by Eq. (23) predicts a general pattern of degeneracies of the $TT$ excitations in linear chains with $q_S = 2, 3, 4, 5, 6, 7, \ldots$ namely $1, 1, 1, 2, 2, 3, 3, \ldots$, i.e., the first and second $TT$ singlet excitation is nondegenerate, the third and fourth excitation is doubly degenerate, etc. In particular, as noted above, the two $TT$ states with $q_S = 4$, the $3^1A_{g}^-$ state, and the $4^1A_{g}^-$ state, should be degenerate in the infinite chain. In a polyene with $N$ carbon atoms the $N/2$ covalent triplets can be combined to a total of $(N^2+2N)/8$ $TT$ excitations. In the limit of large $N$ the degeneracy pattern can be expressed by the equation

$$\rho(q) = \left\{ \begin{array}{ll}
\left(\frac{1}{2}\right)(q - \frac{1}{2}), & \text{if } 0 \leq q \leq N/2 + 1, \\
\left(\frac{1}{2}\right)[N + 1 - (q - \frac{1}{2})], & \text{if } N/2 + 1 < q \leq N + 1.
\end{array} \right.$$  \hspace{1cm} (25)

When summed over $q$, $\rho(q)$ yields the correct total number of $TT$ excitations.

At this point we would like to briefly comment on the question whether the approximate dispersion relations (21) and (22) are asymptotically correct in the limit $k \rightarrow 0$. For the nonalternating polyene with vanishing gaps of covalent triplet and singlet excitations a behavior $E^{-}(k) \sim k$ is strongly supported not only by the corresponding behavior of the spin-wave excitations in the Hubbard model but also by the linear dispersion relations of such excitations that result in the much simpler quantum-mechanical and semiclassical Heisenberg models of the antiferromagnetic chain.\textsuperscript{57} If bond alternation is introduced into the latter model the linear behavior of $E(k)$ remains unchanged, the group velocity of the spin waves decreases like in the PPP model, but, contrary to the latter model, no gap is found.\textsuperscript{54} In contrast, a gap appears to exist in the quantum-mechanical Heisenberg model of the alternating antiferromagnetic chain.\textsuperscript{58} It may well be that the development of covalent gaps in alternating polyenes entails an asymptotic $E^{-}(k) \sim k^2$ behavior for the spin-wave excitations. The fact, however, that Eq. (20) provides accurate excitation energies (see Fig. 10) indicates that such behavior should be confined to small values of $k$ or large values of $N$, i.e., larger than 100.

D. Comparison to spectroscopic observations

The value of 2.87 eV for the asymptotic optical gap (cf. Table I) is about 0.7 eV larger than the value obtained from extrapolations of the $1^1B_{uv}^-$ absorption bands observed for unsubstituted polyenes in solution\textsuperscript{5,59} and about 1.1 eV larger than the value observed in solid state (CH)$_2$.\textsuperscript{60} The former difference has to be attributed probably to our assumption of a bond alternation pattern described by a $\delta$ value which is (1) too large, (2) size independent, and (3) homogeneous,\textsuperscript{13} whereas the latter difference has additional contributions due to "solute effects." All three aspects of the assumed model geometry lead to an overestimate of the covalent as well as of the optical gap energy. For instance, in polyenes the bond alternation is known to decrease from a value $\delta = 0.12\ \text{A}$ for the very short $N = 4$ and $N = 6$ compounds\textsuperscript{45,46} to a value $\delta = 0.08\ \text{A}$ in polyacetylene.\textsuperscript{56} Since the covalent gap arises from bond alternation, the 20% overestimate of the bond alternation in our model geometry ($\delta = 0.10\ \text{A}$) entails the prediction that the covalent gap should be smaller than 1.5 eV. Correspondingly one finds for the optical gap, which decreases less with decreasing degree of bond alternation, an upper bound of 2.7 eV if the 20% overestimate of the bond alternation is taken into account. Solvent effects, which are not accounted for in our calculations and which should lead to a stabilization of the optical state particularly in long chains, reduce the true gap further.

These gap energies predicted for the covalent and ionic bands apply to "vertical" transitions from the ground state, i.e., do not account for vibrational coupling. Because of this latter coupling the so-called "0-0" transitions from the vibrational and electronic ground state to the excited states in their respective vibrational ground state ac-
ually occur at lower energy. Knowledge of the respective energy decrement in finite polyenes\(^{13}\) allows to conclude that the covalent transition in (CH)\(_x\) should start at about 1.3 eV, i.e., in the solid state about 0.5 eV below the optical gap at 1.8 eV, transitions from the ground state extending to higher energies, transitions to the ground state extending to lower energies. Thus, the covalent excitations can explain the long-lived, low-temperature infrared photoluminescence observed at 1.2 eV.\(^{59,60}\) Furthermore, rapid internal conversion from the optical band to the covalent band and subsequent photoexcitation back to the optical band, which has a large oscillator strength, can explain the fast transient photoinduced absorption observed recently at \(\approx 0.45\) eV.\(^{61}\)

E. Effect of dielectric shielding

We have pointed out in our discussion above that electronic excitations in polyenes are affected by the Coulomb interaction between electrons. In realistic systems this interaction can be modified by dielectric screening. We have investigated the effect of such screening and replaced the matrix element describing the Coulomb repulsion (3) by Eq. (4) which, at large distances, corresponds to a dielectric medium with an optical density \(n^2 = 2\). We compare the spectra of unscreened and screened polyenes in Fig. 12(a) (polyenes with bond alternation) and in Fig. 12(b) (polyenes without bond alternation). The results indicate that the asymptotic gaps of the primary covalent and ionic transitions, provided also in Table I, are only slightly affected by the screening. The most important effect might be that the screening lowers the covalent gap in alternating polyenes by about 0.2 eV. In contrast, the influence of the screening on the slopes \(\alpha\) is rather large. Thus, particularly the group velocity of the particle-hole excitations, which is proportional to the slope, increases strongly with increasing screening whereas that of the spin waves decreases only a little.

VI. BOND STRUCTURE AND SOLITONIC EXCITATIONS

Trans-(CH)\(_x\) is known to assume a geometry of alternating bonds.\(^{56}\) At the first glance the ground state of this molecule seems to exhibit two energetically degenerate structures (A) and (B) of alternating double and single bonds, one of which (A) is presented in Fig. 13 (top). The other structure (B) is obtained upon interchanging in (A) double and single bonds. However, a degeneracy of the (A) and (B) phases should exist only in truly infinite (CH)\(_x\) chains. In all finite, i.e., in all real, (CH)\(_x\) molecules which are composed of an even number \(N\) of carbon atoms the ground state is represented by that structure (A) which features the maximum number \(N/2\) of double bonds. The removal of one of these bonds upon replacing, for instance, the singlet spin pair in an ethylene unit by a triplet spin pair requires, as we have shown in the preceding section, an excitation energy \(E^{-}(O)/2 \approx 0.9\) eV. Hence, due to the finite size the degeneracy of structures (A) and (B) is lifted in each (CH)\(_x\) molecule. Structure (B) can be created in part of a (CH)\(_x\) compound from (A) only upon creating a topological soliton-antisoliton pair. A single topological soliton like the neutral soliton depicted schematically in Fig. 13 (bottom) is a domain wall separating the (A) and (B) phases. The ground state of (CH)\(_x\) compounds with an odd number of carbon atoms should have the structure of a neutral soliton. Topological solitons have been inferred to explain a variety of physical properties of

FIG. 12. The effect of dielectric shielding of the Coulomb interaction on the excitation energies of the \(1^1B^+\) and \(2^1A^+\) states of the polyenes for (a) \(\delta = 0.10\) Å and (b) \(\delta = 0.0\). The figure compares the linear \(1/(N+1)\) extrapolations of the lowest covalent (dashed lines) and ionic (solid lines) singlet state excitation energies for the cases of an unsheilded \(U(r)\) and shielded \(U(2r)\) Coulomb interaction.

FIG. 13. One of the two degenerate bond alternation patterns of the ground state of trans-(CH)\(_x\) (top). Schematic representation of a neutral topological soliton in trans-(CH)\(_x\) (bottom).
(CH)_x, in particular its large conductivity upon doping. In this section we want to discuss the relation of the covalent 2^1 A_g^- state to soliton-antisoliton excitations.

Since in our calculations we do not explicitly consider the coupling of nuclear and electronic degrees of freedom we have to resort to empirical relations between \( \pi \)-bond orders and bond lengths like that given by Eq. (6) in order to retrieve information about the polyene geometry in the various polyene states. The resulting values for the order parameter \( \Delta n \) [cf. Eq. (7)] are shown in Figs. 14(a)–14(c) for the 1^1 A_g^+, 1^1 B_u^+, and 2^1 A_g^- states of the polyenes with \( N = 8, 10, \ldots, 16 \) carbons. Solid and dashed lines refer to the results of MRD-CI calculations for alternating \( \delta = 0.1 \) Å and nonalternating \( \delta = 0 \) PPP polyene models, respectively.

Figure 14(a) demonstrates that the order parameter \( \Delta n \) for the ground state assumes uniform positive values corresponding to that geometry of alternating double and single bonds which exhibits a maximum number of double bonds. The predicted alternation of the bond length decreases slightly towards the center of the polyenes, an effect which is more pronounced for longer compounds. For the polyene with \( N = 14 \) carbons the average bond alternation obtained is \( \delta = 0.066 \) Å. The small difference between the two \( \delta \) values demonstrates that the predictions obtained from the empirical bond-order–bond-length relation (6) depend only weakly on the geometry assumed in calculating the electronic ground state. Furthermore, both \( \delta \) values are close to the degree of bond alternation \( \delta = 0.08 \) Å observed in long polyenes. Hence, the bond alternation of the ground state of polyenes is mainly due to the forces exerted by the correlated electron system on the lattice whereas the reverse influence of the lattice on the electrons is smaller. The latter coupling increases the degree of bond alternation by about 40% as measured by the parameter \( \delta \). This result is in harmony with the recent proof that the bond alternation in (CH)_x is mainly due to electron correlation and that the contribution to the alternation caused by the pure (single) electron-phonon coupling, the so-called "Peierls distortion," is of minor importance.

In the excited 1^1 B_u^+ state the order parameter \( \Delta n \) remains positive but, compared to the ground state, decreases towards the central region of the polyenes, corresponding to a weakening of the bond alternation. This behavior is illustrated in Fig. 14(b). For the polyene with \( N = 14 \) carbons the average bond alternation obtained is \( \delta = 0.028 \) Å. The difference between this value and the ground state value \( \delta = 0.08 \) Å explains the large intensity of the C—C single and double bond stretching modes observed in resonance Raman experiments.

The bonding pattern in the excited 2^1 A_g^- state differs from that in the 1^1 A_g^- and the 1^1 B_u^- states. As shown in Fig. 14(c) the order parameter \( \Delta n \) in the 2^1 A_g^- state becomes negative in the central region of the polyenes. Negative values of \( \Delta n \) indicate a reversal of the bond alternation pattern. A change of the sign of \( \Delta n \) corresponds to a soliton structure like that depicted in Fig. 13 (bottom). The order parameter of the 2^1 A_g^- state of \( N = 10, \ldots, 16 \) polyenes shown in Fig. 14(c) character-

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**FIG. 14.** (a) Order parameter \( \Delta n \) of the ground state of the polyenes containing \( N = 8, 10, \ldots, 16 \) \( \pi \) electrons evaluated according to Eq. (7) from PPP-MRD-CI bond orders. The solid lines refer to alternating \( \delta = 0.1 \) Å and the dashed lines to nonalternating model geometries; (b) order parameter \( \Delta n \) of the lowest ionic excited singlet of the \( N = 8, 10, \ldots, 16 \) polyenes; (c) order parameter \( \Delta n \) of the lowest covalent excited singlet state of the \( N = 8, 10, \ldots, 16 \) polyenes.
izes this state as a soliton-antisoliton pair state (see "note added"). The distance between the two solitons measures eight C—C bond units. In case that we assume in our calculation a nonalternating polyene geometry, the soliton-antisoliton structure remains, however, the distance between soliton and antisoliton becomes size-dependent measuring 9 C—C bond units for \( N = 12 \) and 10 C—C bond units for \( N = 14 \). Since in the process of the electronic excitation of the \( 2^{1} A_g^- \) state the \((CH)\_x\) geometry relaxes from an initially alternating structure to a structure with a smaller and partially inverted bond alternation, the increasing distance of the two solitons calculated for the nonalternating polyene appears to indicate that a soliton-antisoliton pair in a long polyene may be formed initially at a small distance and may then oscillate or perhaps separate.

The bond structures of the ground state \( 1^{1} A_g^- \) and the primary covalent excited state \( 2^{1} A_g^- \) can be explained in a simple and illuminating manner. This explanation proceeds along the same lines of reasoning as that on the covalent gap in connection with Fig. 11 and is based on the fact that a double bond in a polyene corresponds to a pair of \( \pi \) electrons at adjacent carbon atoms the spins of which are coupled to a singlet state. The ground state \( 1^{1} A_g^- \) realizes a maximum number of such spin pairings which define the ethylene units of \((CH)\_x\), whereas the \( 2^{1} A_g^- \) state is characterized as involving two intraethylene triple excitations which are coupled to an overall singlet state. The corresponding spin pairings and their relationship to bond structure are illustrated in Fig. 15. In order to arrive at a bond pattern for the \( 2^{1} A_g^- \) state one has to keep in mind that the intraethylene triplets in Fig. 15(b) are coupled to an overall singlet state. This implies that the spins can recouple into a new set of singlet pairings as indicated in Fig. 15(b) by the corresponding connections of pairs of spins. The pairing of spins shown corresponds to the neutral soliton-antisoliton structure also presented in Fig. 15(b). Hence, the singlet spin pairing together with the fact that the primary covalent excitation involves a triplet-triplet excitation with an overall singlet coupling yields the soliton-antisoliton structure of this excitation. The question arises if this composite character of the covalent excitation, i.e., that of a triplet-triplet excitation or that of a soliton-antisoliton excitation, can manifest itself in polyacetylene through an observation of its separate components: Does a covalent singlet excitation yield separate triplet excitations? Does the covalent excitation yield separate solitons?

VII. DISSOCIATION OF COVALENT EXCITATIONS INTO TRIPLETS AND SOLITONS

We have shown in the preceding section that the covalent excitations in polyenes can be regarded either as soliton-antisoliton (SS) or as triplet-triplet (TT) excitations. The wave functions which result from our MRD-CI calculations on finite polyenes describe delocalized excitations, i.e., excitations spread over the entire polyene chain. One expects, however, that for very long polyenes wave packets of these delocalized states can form which are localized in parts of a polyene. The origin of such localization can be either specific interactions exerted when the excitation is initially prepared or can be the coupling to lattice distortions. In the following we want to investigate the latter possibility. If such localization can be achieved it might also be possible to observe the single components of covalent excitations, namely either solitonic excitations or triplet excitations. An argument in favor of this possibility is that the binding energy between two triplets in a TT-excitation is small as shown by the fact that the excitation energy of a TT state is approximately the sum of the excitation energies of two separate triplet states.

The prerequisite that electron-lattice interactions of a magnitude \( \delta E \) can mix pure electronic states \( |j\rangle \) to form wavepackets like

\[
|\omega\rangle = \sum_j a_j |j\rangle
\]

is that the energy separation between the states \( |j\rangle \) does not exceed \( \delta E \) by a large margin. Furthermore, our results on the spectra of polyenes as shown in Fig. 10 allow an estimate of the level density of covalent states. As we have demonstrated in Sec. V the low-lying covalent states in alternating polyenes with \( \delta = 0.10 \) Å approximately obey the simple behavior

\[
E_q(N) = \Delta E_0 + q\frac{\alpha}{N+1}, \quad q = 2, 3, 4, \ldots
\]

where the numerical constants, given in Table I, are determined by the parametrization of the PPP Hamiltonian (2). In the limit of large \( N \) the degeneracy \( \rho(q) \) of these states
is given by Eq. (25) such that we can estimate the density of covalent states in long polyenes in the energy range above $\Delta E_0$.

With respect to their coupling to the lattice there should be a qualitative difference between dissociated $SS$ and $TT$ excitations in long polyenes. Dissociated $SS$ excitations are topological excitations that require the distortion of the complete lattice between soliton and antisoliton whereas dissociated $TT$ excitations require strictly local distortions only. This difference is schematically indicated by the difference between the bond structures ($\alpha$) in Figs. 15(b) for the $SS$-pairs and 11(b) for the $TT$ excitations. An upper bound for the electron-lattice interaction energy $\delta E_0$, estimated from Eq. (5) as the difference of the resonance integral $t$ for single and for double bonds, is $0.34$ eV. This upper bound for $\delta E_0$ should apply to the case of a topological lattice distortion connected with a distant $SS$ pair. However, in the case of a strictly local lattice distortion delocalized states will only experience an effective interaction of about $0.34$ eV/N since local amplitudes of delocalized states are proportional to $N^{-1/2}$. The question arises if an interaction of that order of magnitude suffices to induce localized covalent excitations in long polyenes, i.e., two localized triplets and close $SS$ pairs. Since the spacing between covalent states according to Eq. (27) is also proportional to $1/N$, perturbation theory does not provide an answer. Instead we need to derive a nonperturbative estimate. For this purpose we introduce the following strong assumption regarding the matrix elements of the perturbation $\delta V$ originating from a local distortion of the lattice

$$\langle j | \delta V | k \rangle = \delta E < 0,$$

i.e., the values of these matrix elements are assumed to be independent of $|j\rangle$ and $|k\rangle$ and negative. Here $|j\rangle$ and $|k\rangle$ are delocalized covalent excitations as defined in (26). The latter assumption is plausible for both $SS$ and $TT$ pairs because from the $\pi$-bond orders of the covalent states $|j\rangle$ one expects $\delta E$ to be negative if the distorted lattice exhibits a less distinct bond alternation than that defined by $\delta = 0.10$ Å. Such lattice distortion, however, should accompany localized triplets as well as solitonic excitations. The eigenvalue problem of the perturbed band of covalent excitations is defined by the equation

$$\epsilon j |a_j\rangle + \delta E \sum_k a_k = \epsilon a_j ,$$

where $\epsilon j$ and $a_j$ are the energies and expansion coefficients of the unperturbed states $|j\rangle$. In the following all energies will be defined relative to the band edge $\Delta E_0$. The expansion coefficients can be expressed formally as follows

$$a_j = \frac{\delta E}{\epsilon - \epsilon_j} \sum_k a_k .$$

Iterating this equation once yields

$$1 = \delta E \sum_j \frac{1}{\epsilon - \epsilon_j} .$$

The solution of Eq. (31) provides the energy spectrum $\epsilon j$ of the perturbed band of covalent states. This spectrum is located close to the unperturbed band of energies $\epsilon_j$ except for the eigenvalue of one state which is moved far below the unperturbed band. To demonstrate the existence of this state we consider Eq. (31) in the limit of large $N$. Then one can replace the sum by an integral to obtain

$$1 = \delta E \int_0^{N+1} \frac{\rho(q)dq}{\epsilon - \epsilon_q} ,$$

where the energies $\epsilon_q$ of the covalent states are given by Eq. (27) and their degeneracies $\rho(q)$ by Eq. (25). The integral can be evaluated and in the limit $|\epsilon| \gg \alpha$ one finds

$$\epsilon \approx \frac{\delta E}{4} N^2 \approx -(0.09 \text{ eV}) N .$$

In fact, this expression proves that the interactions in the band of covalent excitations induce a state which for large $N$ is strongly shifted and, hence, localized.

The important property of relationship (33) is that the energy shift increases linearly with $N$, i.e., one expects that for some length $N$ of a polyene localization will set in. However, it is impossible to estimate this length since expression (33) overestimates the energy shift. There are three reasons. First, the assumption $|\epsilon| \gg \alpha$ used in the derivation entails a crude approximation of the energy $\epsilon$ of the localized state. Second $\delta E_0 \approx 0.34$ eV/N only provides an upper bound for the coupling to lattice distortions since we assumed an extreme reversal of single and double bonds in estimating its value. Third, the matrix elements of the perturbation are not as homogeneous as assumed by Eq. (28) but rather one has to expect a spread of the values of these matrix elements and even sign reversals. All that implies a reduction of the energy shift. Nevertheless, Eq. (33) indicates that if the interaction $\langle j | \delta V | k \rangle$ is homogeneous over a sufficiently broad band of energies a local lattice distortion can produce localized excitations. The possibility that topological distortions like solitons can be formed by coupling to the lattice follows in a straight forward way by perturbation theory.

We want to argue now that there exists indeed strong experimental evidence for a separation of the covalent states in polydiacetylene into separate, i.e., localized triplets. Such triplets would be observable after photoexcitation of the material into the band of ionic states, internal conversion to the band of covalent states and dissociation of a $TT$ state into separate triplets. Evidence for the appearance of fast photoinduced triplets has been provided recently by Robins et al. on the basis of magnetic field modulated optical absorption and microwave absorption experiments. However, the observations of these authors do not reveal the route of triplet production. In a recent microwave absorption experiment with nanosecond time resolution Sixl et al. found that initially all triplets sublevels are equally populated, i.e., the triplets formed are completely unpolarized. Triplets formed through the strongly orientation-dependent interactions of an intersystem crossing route are always polarized to some degree. The puzzling observation of Sixl et al. can be explained readily, however, if the triplets are born from a covalent $TT$ excitation. In this excitation the two triplets denoted
by $T_{1}^{(1)}$ and $T_{2}^{(2)}$ are coupled to an overall singlet state $S_0^*$ according to the scheme

$$S_0^* = \frac{1}{\sqrt{3}} (3T_{1}^{(1)} 3T_{-1}^{(2)} - 3T_{0}^{(1)} 3T_{0}^{(2)} + 3T_{1}^{(1)} 3T_{1}^{(2)}).$$

(34)

Since the numerical (Clebsch-Gordan) coefficients for all triplet sublevels, except for a phase factor, are identical in this scheme neither of the triplets is polarized. This follows formally from the fact that the matrix elements for the spin operator $S_z$ of triplet $T_m^3$, i.e., $\langle S_0^* | S_0^y | S_0^y \rangle$, $S_\beta=x,y,z$, vanish. Consequently, the observation of Sixl et al. proves unequivocally that the covalent singlet excitations are the parent states of the photoinduced triplets observed in polydiacetylene. We note that the proposed $TT$ state dissociation is analogous to the singlet exciton fission observed in organic crystals.65,66

The covalent excitations may also decay into separate spin-1/2 solitons which have been characterized schematically in Fig. 13 (bottom), though no proof exists so far for this process. The solitons formed by this route should give rise to photoinduced ESR signals which also should be unpolarized because of their singlet parent state. The decay route, i.e., either to triplets or to solitons, should depend on the material involved. For example, polydiacetylene's bond structure is unfavorable for soliton formation, hence the triplet route is more favorable. Polyacetylene can accommodate solitons, hence the decay to solitons should be observed in case the decay to triplets is less efficient. This might actually be the case since the perturbation for triplet localization is $BE \sim 1/N$, whereas that for solitons is likely to be independent of $N$.

The fact that photoinduced triplet pairs and possibly also soliton pairs are born in a pure singlet quantum state can be proven by a magnetic field dependence of their recombination dynamics. If a large enough fraction of triplet pairs or solitonic spin-1/2 pairs in polyacetylene at a time $t$ after their birth can reform to the singlet covalent state or to the singlet ground state this process acts as a filter for the singlet character of the pair at time $t$. An external magnetic field acting together with either the fine structure interaction (triplet) or the hyperfine interaction (spin-1/2 solitons) leads to a yield of free triplets or solitons with a characteristic magnetic field dependence. The theory of this process has been described for such recombination processes in liquids in Ref. 67.

VIII. SUMMARY AND OUTLOOK

The properties of polyacetylene and of related one-dimensional conducting and semiconducting polymers are dominated by the effects of electron correlation. The polymers are not of "infinite" length but typically consist of a few hundred to a few thousand elementary units. The description of the excitations of a correlated electron system that is neither "infinite" nor "small" falls into a domain of physics that is located between molecular physical and solid state physics. In this paper we have demonstrated, using polyacetylene as an example, how these branches of physics can be bridged. Our approach is based on four fundamental assumptions the validity of which has been investigated in detail.

(1) The physical nature of excitations in infinite and in large, but finite, many-electron systems is identical. Thus, the energy-momentum relations valid in the infinite system for the various classes of excitations, like covalent (spin-wave) or ionic (particle-hole) excitations, should apply also to finite systems.

(2) In finite systems the wavelength of an excitation cannot exceed the system length $L$ if the system is cyclic and $2L$ if it is linear. According to the de Broglie relation, therefore, the energetically lowest excitations of each class is associated with an elementary transition momentum. Equations (9) and (10) give the lower bounds for the elementary transition momenta in cyclic ($N=4n+2$) and linear $N$-electron systems, respectively.

(3) The values of the elementary transition momenta are solely determined by the geometry of the finite system, i.e., are given by the lower bounds.

(4) The momenta associated with transitions to higher excited states are integer multiples of the elementary transition momenta [cf. Eqs. (11) and (12)]. Consequently, as expressed by Eqs. (14) and (15), the discrete energies of a finite system are obtained by evaluating the energy-momentum relations of the infinite system at multiples of the elementary transition momenta. Conversely, the dispersion relations of the infinite system can be determined from the energy spectra of finite systems if the values of the transition momenta are known.

For the investigation of the validity of these assumptions we have first selected the Hubbard model since it is the most simple model of one-dimensional many-electron systems that describes both the kinetic energy and the Coulomb interaction of the electrons, and since it is the only model for which exact results on finite as well as on infinite systems are available. In those cases, in which exact excitation energies for finite systems were unavailable, we have employed a multireference double-excitation configuration interaction expansion (MRD-CI) which recently has been developed by us35 for the calculation of accurate excitation energies. We have demonstrated that assumptions (1), (2), and (4) render excellent and asymptotically (for large $N$) correct approximations. Assumption (3) has been shown to be valid for covalent (spin-wave) excitations in rings and chains and for ionic (particle-hole) excitations in rings, whereas it turned out to be invalid for ionic excitations in chains. The transition momentum associated with the lowest ionic excitation in chains was found to increase with increasing Coulomb interaction of the electrons [see Eq. (13)]. Such increase, though possibly smaller, appeared to exist also for higher ionic excitations. A more detailed study of these effects requires analytical results on the Hubbard model of finite linear chains. It would be desirable, in this respect, to derive equations for finite Hubbard chains that are analogous to the Lieb-Wu equations33 for finite rings.

For Hubbard interaction parameters, which apply to $\pi$ electrons in polyenes, the dispersion relations of ionic and covalent excitations are nearly linear over a wide range of momenta. In this linear regime assumptions (1)–(4) allow
the construction of approximate dispersion relations from the energy spectra of finite polyenes by means of $1/(N+1)$ extrapolations. Assuming that a linear approximation of the energy-momentum relation also holds for polyenes described by the Pariser-Parr-Pople (PPP) model we have evaluated dispersion relations for nonalternating and alternating polyenes using the results of MRD-CI calculations.

The gap of the band of covalent singlet excitations is found to vanish for nonalternating polyenes and to be large for alternating polyenes. We have estimated for this gap in (CH)$_2$ a value of about 1.3 eV.

The covalent singlet states of long polyenes can be characterized as being composed of two or more triplet spin waves combined to an overall singlet state. This characterization allowed the derivation of the momentum quantum numbers of the covalent singlet states from those of the triplet states and of the state density in the covalent band. The same connection between covalent triplet and singlet states has been exploited by Woynarovich for the Hubbard model in his derivation of singlet spin-wave dispersion relations from those of triplet states.\footnote{W. P. Su, J.-R. Schrieffer, and A. J. Heeger, Phys. Rev. Lett. 42, 1698 (1979).}

The dispersion relations of the covalent states in nonalternating polyenes are strictly linear in the limit of small momenta. It is uncertain if that is also the case for alternating polyenes which exhibits a covalent gap. The compounds investigated by us are not long enough to allow definitive statements. For an answer to this question one should try to obtain an analytical solution of the Hubbard model of alternating polyenes (using periodic boundary conditions).

For the ionic excitations only the asymptotic "optical" gap has been estimated from extrapolations. No dispersion relations could be derived because in linear chains the transition momenta depend on the Coulomb interaction of the $\pi$ electrons and can be determined from extrapolations only up to a scaling factor. Due to electron correlation the optical gap is large ($\approx 2.2$ eV) in nonalternating polyenes and is found to increase by about 40% upon bond alternation.

The bond orders calculated from the MRD-CI wave functions have provided insight into the bond structure of long polyenes in ground and excited states. It has been found that covalent singlet states exhibit the bond structure of a soliton-antisoliton pair. The density of states in the covalent band has been estimated to be high enough to allow in long polyenes a localization of such excitations through coupling to the lattice degrees of freedom. We have interpreted the results of recent microwave experiments\footnote{S. G. Kivelson and D. E. Heim, Phys. Rev. B 26, 4278 (1982).} as evidence for the decay of covalent singlet states into separate localized triplets.

Note added. Recently Hayden and Mele have presented results of renormalization-group (RG) calculations on a 16-site Hubbard-Peierls chain.\footnote{P. Horsch, Phys. Rev. B 24, 7351 (1981).} The order parameters calculated for the ground and $1 B_{2u}^+$ states are similar to those shown in Fig. 14(a) and 14(b). But for the $2 A_{1g}^-$ states a structure of four solitons is predicted instead of the soliton-antisoliton structure exhibited in Fig. 14(c). The prediction of an additional soliton-antisoliton pair, located at the center of the 16-site chain, may be an artifact of the RG procedure that constructs 16-site wave functions from two 8-site chains employing quite restricted many-electron basis sets.

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