Prospects for continuous-wave molecular modulation in Raman-active microresonators

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Abstract: We propose the extension of existing molecular modulation techniques for continuous-wave optical modulation to Raman-active microresonators. Intense pump and Stokes modes inside a microresonator prepare a high coherence of the Raman transition between two ro-vibrational states. With the coherence prepared, any laser in the optical region can be coupled to the microresonator and be modulated. We perform numerical simulations which predict that these “microresonator-based molecular modulators” could have modulation efficiencies on the order of 1% at 10 THz-scale frequencies for any optical wavelength.

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1. Introduction

Optical modulators are powerful tools that have found a wide variety of uses, ranging from encoding information to synthesizing ultrashort waveforms. Acousto-optical and electro-optical modulators (AOMs and EOMs) provide frequency shifts limited to the electrical frequencies that drive them, with a maximum of a few hundred GHz. Although new techniques continue to incrementally improve electrically-driven modulators [1–5], there is little prospect for pushing them to significantly higher modulation frequencies. Optical modulators with THz-scale modulation frequencies need to be driven by optical processes.

Over the past decade our group created a 90 THz continuous-wave (CW) optical modulator based on a technique called molecular modulation, a kind of driven four-wave mixing based on CW Raman lasing [6–15]. Molecular modulation was originally developed using pulsed pump lasers [16–27], but these systems suffer from large linewidths and low repetition rates [28]. Using CW driving beams for our modulator avoided these problems, but required the use of an optical cavity to reach the high intensities necessary for molecular modulation to occur. Using a large gas-filled cavity ultimately limited the usefulness of the modulator due to its complexity of operation and sensitivity to mechanical and thermal fluctuations, all of which negatively impacted the steady-state modulation efficiency $\epsilon$.

In this work we propose the extension of CW molecular modulation to solid-state Raman-active microresonators and discuss the prospective characteristics of such a device, which we call a Microresonator-based Molecular Modulator (MMM). We also present numerical simulations that indicate that an MMM based on a silica microsphere could achieve a frequency shift of $\sim 12$ THz with modulation efficiencies of $\epsilon \approx 1\%$ at any optical wavelength.

Microresonator modes have small volumes and high quality factors over a wide range of wavelengths, which make them ideal platforms for building nonlinear optical devices [29,30]. For example, Grudinin et al. demonstrated an ultra-low-threshold Raman laser in a CaF$_2$ microresonator which required just 15 µW of launched pump power [31,32]. Other groups have used microresonators to create high-quality stabilized optical frequency combs or explore more exotic nonlinear behavior such as soliton formation [33–35]. As an added benefit, the smaller overall system size and better mechanical and thermal stability offered by microresonators will help mitigate the issues that limited the efficiency of our previous device. Microresonators could also be integrated into complex optical systems much more readily.
Figure 1 shows a cartoon of the specific kind of proposed MMM we characterize in this work. A pump laser is evanescently coupled to one of the whispering gallery modes of a silica microsphere using a tapered fiber [36]. This “pump mode” builds up to high intensity. The pump mode then generates a high-intensity Stokes mode through stimulated Raman scattering (SRS). Once the pump and Stokes modes have both built up to high intensities, molecular modulation can occur: a separate low-power “mixing” beam is coupled to the resonator and is modulated via driven four-wave mixing to produce light at the “target” frequency.

Fig. 1. A cartoon of our molecular modulation scheme implemented using a microsphere made of Raman-active material with frequency shift $\omega_m$ coupled to a tapered fiber. The launched pump beam at frequency $\omega_P$ drives a Stokes mode at frequency $\omega_S = \omega_P - \omega_m$ via Raman lasing. Then a launched “mixing” beam at frequency $\omega_M$ is modulated by the intense pump and Stokes modes via four-wave mixing to produce light at the “target” frequency $\omega_T = \omega_M + \omega_m$.

2. Theoretical modeling

To characterize the theoretical performance of MMMs we have derived a model of the four-wave mixing process in an arbitrary resonator. Typically, models of this process either do not perform a spatial decomposition at all (leading to the “sideband model”) [37,38], or assume the well-known regular mode structure of a Fabry-Perot cavity [7,39,40]. In contrast, we decompose the electric field into the spatial modes of a generalized resonator as well as in time. We combine this description with models of the underlying quantum mechanics and of how light couples to the resonator (see [36,41,42]) to create an end-to-end, time-dependent model of the modulator.

The primary advantage of this model is that it makes no assumptions about the mode structure of the resonator or which inter-mode couplings might be more important than others. Instead, these relationships will arise emergently from the model, as they would in a real experiment. As we will see, this turns out to be critically important to understanding the behavior of the MMM.

An additional advantage of our model is that it includes all of the time-dependent behavior of the resonator (i.e., it is not solely concerned with steady-state behavior). We can use this to look at, for example, how long it takes the mode energies to reach steady-state after the pump laser turns on. Knowing that timescale will be critical for building optical modulators that can interoperate with other devices. A delay of several microseconds before the output power stabilizes could be important for both downstream and upstream devices. We plan to explore this aspect of the model in future work.

The full derivation of the model is too long to be included here, but we will briefly sketch it. Instead of starting from a set of classical master equations, we begin with a low-level description of the underlying quantum mechanics, to ensure that we capture the physics we are looking for (as expected, the final result is equivalent to what one would get by starting with the classical description). We begin with a decomposition of the molecular state as a function of position in the cavity $r$ and time $t$ into two ro-vibrational states $|a\rangle$ and $|b\rangle$ with frequencies $\omega_a$ and $\omega_b$.
where $u$ (frequencies $\omega_i$) [40]:

$$|\Psi(r,t)\rangle = c_a(r,t) e^{-i\omega_st} |a\rangle + c_b(r,t) e^{-i\omega_st} |b\rangle + \sum_i c_i(r,t) e^{-i\omega_it} |i\rangle .$$  

(1)

The Hamiltonian in this basis is $\hat{H} = \hat{H}_0 + \hat{H}_f$, where

$$\hat{H}_0 = \hbar \left[ \omega_a |a\rangle\langle a| + \omega_b |b\rangle\langle b| + \sum_i \omega_i |i\rangle\langle i| \right]$$  

(2)

$$\hat{H}_f = -\sum_i \mathcal{E}(r,t) \cdot \mu_{ai} |a\rangle\langle i| - \sum_i \mathcal{E}(r,t) \cdot \mu_{bi} |b\rangle\langle i|$$

$$-\sum_i \mathcal{E}^*(r,t) \cdot \mu_{ai}^* |i\rangle\langle a| - \sum_i \mathcal{E}^*(r,t) \cdot \mu_{bi}^* |i\rangle\langle b|$$  

(3)

and where $\mathcal{E}$ is the electric field vector and $\mu_{xy}$ is the dipole moment between states $|x\rangle$ and $|y\rangle$. There are no direct transitions between states $|a\rangle$ and $|b\rangle$ (i.e., $\mu_{ab} = 0$).

We decompose the electric field into cavity modes indexed by $q$ (frequencies $\omega_q$).

$$\mathcal{E}(r,t) = \frac{1}{2} \sum_q [\mathcal{E}_q(t) e^{-i\omega_q t} \mathbf{u}_q(r) + \mathcal{E}_q^*(t) e^{i\omega_q t} \mathbf{u}_q^*(r)].$$

(4)

where $\mathbf{u}_q(r)$ is the mode shape function for mode $q$. We use lower-case subscripts for generic mode indices and upper-case color-coded subscripts for specific modes throughout. An energy level diagram for this system is shown in Fig. 2.

![Energy Level Diagram](image)

**Fig. 2.** The energy level diagram for molecular modulation. Two stable molecular ro-vibrational states $|a\rangle$ and $|b\rangle$ interact with four photons via two intermediate virtual states (both labeled $|i\rangle$). The Raman frequency shift $\omega_m$ is the frequency difference between the molecular states. The arrows are in the correct direction for term $\text{MSP} \rightarrow \text{T}$.

Armed with this decomposition, we eliminate the electronic states $|i\rangle$ from the evolution equations for $c_a(r,t)$ and $c_b(r,t)$, assume that the interaction is perturbative, and solve for the molecular polarization in terms of the $\mathcal{E}_q$. This polarization is inserted into the evolution equations for the $\mathcal{E}_q$ themselves, which include additional terms for the intrinsic decay of the mode energy as well as input power from launched beams coupled to the resonator [41,42]. After much simplification, we arrive at a final result that is similar in form to the sideband model:

$$\dot{\mathcal{E}}_q(t) = -\frac{\omega_q}{2Q_q} \mathcal{E}_q + \frac{1}{2} \sqrt{\frac{\omega_q}{Q_q}} \sum_i \sqrt{\delta_i} e^{-i(\omega_q - \omega_i)t}$$

$$+ i \sum_{r,s,t} G_{rstq} \omega_q \mathcal{E}_r \mathcal{E}_s^* \mathcal{E}_t \frac{V_{rstq}}{V_{qq}} \left( \frac{1}{\delta_{s,t}} + \frac{1}{\delta_{s,t}} \right) e^{i\omega_{rst}t},$$

(5)
where \( q \) is the index of the cavity mode whose derivative we are evaluating, and \( r, s, \) and \( t \) are indices for a sum over all of the cavity modes (including mode \( q \)).

Each mode \( q \) has an intrinsic quality factor \( Q_q^r \), which describes the decay of the mode’s stored energy, and a coupling quality factor \( Q_q^c \), which describes how easily light can move from it into the fiber or vice-versa. The total effective quality factor for each mode \( Q_q \) is given by \( 1/Q_q = 1/Q_q^r + 1/Q_q^c \). The first term of Eq. (5) thereby accounts for both the intrinsic decay of the mode energy and the power that couples out to the fiber.

The second term is the power that couples into the mode from the fiber. It keeps track of each launched beam \( l \) with frequency \( \omega_l \) and power \( S_l \) independently. The factor \( E_q \) is the prefactor in the mode energy \( U_q = E_q |E_q|^2 \).

The terms in the sum are the nonlinear four-wave mixing interaction, with a mode-dependent Raman gain coefficient \( G_{rstq} \). Two volume factors appear; both are volume integrals of mode shapes \( u_q \) over the resonator volume, defined by

\[
V_{abc...} = \int d^3r |u_a||u_b||u_c|\cdots.
\]

\( V_{rstq} \) is a four-mode shape overlap, and \( V_{qq} \) is simply the mode volume. Two frequency differences appear in these terms: the two-mode Raman detuning \( \delta_{rs} = (\omega_r - \omega_s) - \omega_q + \gamma_R \) (where \( \gamma_R \) is the Raman linewidth), and the four-mode detuning \( \Delta_{rstq} = \omega_r - \omega_s + \omega_t - \omega_q \). For convenience, we consolidate the various prefactors of each term to get the following form:

\[
\hat{E}_q = -D_q E_q + P_q \sum_{l} \sqrt{S_l} e^{-i(\omega_q - \omega_l)t} + \sum_{r,s,t} G_{rstq} E_r E_s E_t e^{i\Delta_{rstq}t}.
\]

The primary differences between Eq. (7) and the sideband model are that we have not restricted the sum over possible four-wave mixing terms, and that we have allowed more than one launched beam. One can reduce Eq. (7) to the sideband model by restricting the sums appropriately, and we have confirmed that in that form the model reproduces expected behavior such as clamping during cascaded SRS [38], which we also see in the full model.

Even in the fully-general case, the most important terms in the sum for a given mode \( q \) can be identified in advance by a simple graphical algorithm using the energy level diagram in Fig. 2. We label the terms by the four modes they involve, so the term \( i\mathcal{G}_{MMPT} E_M E_P E_T \exp(i\Delta_{MMPT}t) \) is labeled \( \text{MSP} \rightarrow T \). The notation indicates that this term is the one where modes \( M, S, \) and \( P \) contribute to the derivative of \( T \).

The arrows on the diagram must form a closed loop (their directions may need to be reversed) to have small \( \Delta_{rstq} \) (see Sec. 3.2), so important terms must be either of the form \( qrr \rightarrow q \) (two pairs of modes) or \( rst \rightarrow q \) (all four modes different). Additionally, arrows that do not touch the same energy level must not appear next to each other (e.g., \( T \) cannot appear next to \( S \)). Finally, the last mode’s arrow must point upward (this corresponds to an arbitrary sign convention in the derivation and has no physical significance; if the sign choice were reversed, the term with the last arrow pointing downward would be the important one).

For example, the most relevant terms for the target mode \( T \) must be of the forms \( Trr \rightarrow T \) or \( rst \rightarrow T \). For the first form, \( rr \) could be either \( MM, PP, \) or \( SS \). These correspond to the possibility of stimulated Raman scattering (SRS) between the target and mixing, pump, or Stokes modes. For the second form, the \( T \) arrow must point upward, so we are forced to go around the diagram in an order that gives us the term \( \text{MSP} \rightarrow T \). This term corresponds to molecular modulation: it says that energy can be delivered to the target mode from the other three modes via four-wave mixing.
3. Computational modeling

We can make some predictions about the behavior of an MMM by simulating Eq. (7) numerically. In this section we will first lay out the assumptions behind our simulations, then discuss the dependence of the modulation efficiency $\epsilon$ on a variety of parameters.

We evolve the mode amplitudes in time using a simple Runge-Kutta scheme with a time step of a few picoseconds, which is enough to resolve frequency detunings of up to a few hundred GHz (more than large enough to resolve all of the important terms in Eq. (7)). For consistency, four-wave mixing terms in Eq. (7) that have a $\Delta_{qrst}$ larger than the Nyquist frequency for the chosen time step are not included in the calculation. The simulations are converged with respect to the time step and always reach a steady-state with constant energy in each mode. The final energies do not depend on the relative timing of the launched pump and mixing beams being turned on, so we turn both beams on at the beginning of the simulation.

We take the resonator to be a silica microsphere with radius $R = 50 \mu m$. We model the silica Raman shift as a Lorentzian with $\omega_m = 2\pi \times 12$ THz and a linewidth of 2 THz, and the $G_{rstq}$ are calculated from the bulk Raman coefficient [43]. We take the intrinsic quality factors of all of the modes to be $10^8$ (quality factors this high have been regularly achieved in practice [36]), except in Sec. 3.4 where we look at how the modulation efficiency depends on it.

Light is coupled to and from the resonator using a tapered fiber. We generally either calculate the coupling quality factors from Eq. (35) of [44] with a 1 $\mu m$ taper radius and the pump mode critically coupled to the launched pump beam (“pump-critical”), or we take all of the modes to be critically coupled (“all-critical”). The pump-critical case does not give the highest modulation efficiency, but it does have the lowest threshold (this will be addressed in Sec. 3.5).

The all-critical case is roughly equivalent to adding a second tapered fiber to the system, with coupling optimized for the mixing mode instead of the pump mode. This kind of coupling scheme is common for communications applications [45,46], and could be re-purposed for the MMM. It would be necessary to prevent the secondary fiber from disturbing the coupling between the first fiber and the microsphere, perhaps by suppressing any propagating fiber modes near that wavelength using an engineered stopband [47]. However, the purpose of this case is just to approximate the maximum possible theoretical efficiency of the system, and in our simulations we simply assume it might be possible without implementing a model of how it would work in practice. As we will see in Sec. 3.5, even this is not the most ideal coupling scheme.

For better control over the parameters of the simulation, we use representative modes instead of real cavity modes. The mode volumes and four-mode shape overlaps of the representative modes are given values calculated from averages over real cavity modes, while we set the modes frequencies and intrinsic quality factors arbitrarily. It is important to control the mode frequencies exactly because the modulation efficiency is quite sensitive to the four-mode detuning $\Delta_{qrst}$, which is a measure of the timescale over which the modulation process conserves energy. We will generally set the four-mode detuning for the molecular modulation path to zero. A possible method for control of the four-mode detuning will be discussed in Sec. 3.2.

As mentioned above, the mode volumes $V_{qq}$ and four-mode shape overlaps $V_{rstq}$ are given uniform values for all modes that are representative of the average volumes and overlaps calculated from real cavity modes [36,48]. The actual mode volumes and four-mode shape overlaps vary up and down somewhat-independently from mode to mode over the range of mode wavelengths we use by approximately an order of magnitude, depending on the wavelengths of the modes, their angular momentum and radial quantization numbers, and their polarizations. Since the four-wave mixing terms in Eq. (5) depend only on the ratio between a four-mode overlap and a mode volume, we expect the real modulation efficiencies to be within an order of magnitude of those calculated with the representative values.

For simplicity, we assume that cascaded SRS has been suppressed. This could perhaps be achieved in practice [49], but actually turns out to be irrelevant. We have run the same simulations
with enough higher-order Stokes modes to capture the full cascade at the highest launched pump powers that we show below, and there is no significant difference in the modulation efficiency compared to simulations without the cascaded modes (this surprising result will be discussed in Sec. 3.1). The launched mixing power is always $S_M = 1\,\mu W$, which is below the threshold for the mixing mode to create its own Stokes mode.

Our simulations do not take into account other nonlinear processes, such as generation of parametric sidebands through self-phase modulation, which is due to the Kerr nonlinearity [50]. The Kerr and Raman nonlinearities are both third order, so one would expect their thresholds to be comparable. In a real experiment, there will likely be additional modes excited near the pump mode due to Kerr mixing. However, there is good reason to believe that the existence of these additional excited modes will not significantly change our results.

First, the primary effect of these modes will be to deplete the energy of the pump mode. However, in Raman lasing, the pump energy inside the resonator is clamped once it reaches the Stokes lasing threshold [38]. When a Raman laser is pumped far above threshold, a large fraction of the incident pump power does not couple into the resonator. This excess incident pump power can overcome any additional depletion of the pump beam due to the presence of other nonlinear optical processes. Second, the additional four-wave mixing pathways made available by these extra excitations will not, in general, produce a target frequency that is aligned with a resonator mode frequency (i.e., these terms will have large four-mode detunings). As we will see in Sec. 3.2, this should suppress any additional modulation of the mixing beam. However, a detailed analysis of all of these additional nonlinear processes in addition to Raman generation is beyond the scope of this work, and possibly too computationally expensive to explore with the mode-based simulations we use.

3.1. Dependence on launched pump power

An important metric for a modulator is the modulation efficiency $\epsilon$, which we define to be the output power of the target mode $P_T$ divided by the launched mixing power $S_M$ ($\epsilon = P_T / S_M$). Since the molecular modulation process only relies on establishing coherence between the molecular states through SRS, we should only expect non-zero modulation efficiency as soon as the pump mode energy goes above the threshold to produce a Stokes mode. Figure 3 shows that this is indeed the case.

The modulation efficiency curves are quite unlike what we expected based on the behavior of our previous device. In that system the mixing and target frequencies were not resonant in the cavity, so the target light was produced in a single pass of the mixing beam through the system. The modulation efficiency was very low, so the pump mode, Stokes mode, and mixing beam were essentially unperturbed by the molecular modulation process. This allowed the modulation efficiency to scale roughly like the Stokes mode energy in the ideal case. In contrast, the situation in the MMM is shown in Fig. 4. The mixing mode energy decreases after an initial plateau, corresponding to the decrease in the modulation efficiency.

To investigate this counterintuitive decrease, we start by noting that the complex mixing mode amplitude $\hat{E}_M$ is constant at steady-state (not just the mode energy $U_M$) because it is externally pumped. Therefore, we can set the derivative of $\hat{E}_M$ to zero at steady-state, assume that it is too far detuned from the launched pump beam to interact with it, and write

$$\dot{\hat{E}}_M = 0 = -D_M \hat{E}_M + P_M \sqrt{S_M} + i \sum_{r,s,t} G_{rstM} \hat{E}_r \hat{E}_s^* \hat{E}_t e^{i \Delta_{rstM} t}$$

(8)

$$E_M = \frac{P_M}{D_M} \sqrt{S_M} + i \sum_{r,s,t} \frac{G_{rstM}}{D_M} \hat{E}_r \hat{E}_s^* \hat{E}_t e^{i \Delta_{rstM} t}.$$  

(9)

This equation does not allow us determine $E_M$ from the initial conditions, but it does allow us explain why $E_M$ has the steady-state value it does phenomenologically. The most important terms in
Fig. 3. The modulation efficiency $\epsilon$ as a function of launched pump power $S_P$ for three different mixing wavelengths $\lambda_M$. The modulation efficiency is $\epsilon = P_T / S_M$, where $P_T$ is the target mode output power and $S_M$ is the launched mixing power. The pump mode wavelength is $\lambda_P = 1064$ nm, and it is critically coupled to the launched pump beam.

Fig. 4. The energy stored in each resonator mode $U_q$ as a function of launched pump power $S_P$. After threshold, $U_P$ is clamped and $U_S$ grows like the square root of the launched pump power. After an initial plateau following threshold, $U_M$ and $U_T$ begin to decrease, which reduces the modulation efficiency.

Eq. (9) are shown in Fig. 5 for one of the curves from Fig. 3. Note that all of these terms have a four-mode detuning of zero in these simulations, so they do not have time-dependent phases that do not arise from the mode amplitudes themselves. As expected, the launched mixing beam and the four wave mixing are both important. However, another term is also important:
\(\text{MSS} \rightarrow M\), which corresponds to SRS between the mixing mode and the Stokes modes (the next most important is \(\text{MPP} \rightarrow M\), not shown).

![Graphical representation of mixing mode and Stokes mode interactions](image-url)

**Fig. 5.** The real and imaginary contributions of the three most important terms in Eq. (9) to the mixing mode amplitude (magnitude plotted as \(|E_M|\)): the launched mixing beam, the molecular modulation term \(\text{TPS} \rightarrow M\), and the SRS term \(\text{MSS} \rightarrow M\). These three terms together almost entirely compose the steady-state mixing mode amplitude (the magnitude of their sum is plotted in dotted green).

SRS would normally be suppressed between these modes because they are not on Raman-resonance. However, the Raman linewidth of silica is several THz, so the frequency difference is not enough to suppress this term when the Stokes mode energy is large. Although the simulations we present here only include one Stokes mode, we have performed similar simulations with enough higher-order Stokes modes to model the entire cascade out to \(S_P = 10\ W\), and there is no significant difference in the behavior of the modulator. The combined effect of all of the cascaded Stokes modes together is similar to the single large un-cascaded Stokes mode. This effect is why there is no significant difference in modulation efficiency between simulations run with and without cascaded Stokes modes.

Remarkably, this off-resonant SRS actually supports the mixing mode, causing the plateau. We ran a set of simulations with the Raman linewidth reduced by a factor of \(10^3\) (i.e., it is a few GHz instead of a few THz) while keeping everything else constant. This suppresses the off-resonant SRS between the mixing mode and pump or Stokes modes. The mode energies and mixing mode amplitude composition for these simulations are shown in the left and right panels of Fig. 6 respectively. With no SRS from the mixing mode to the pump or Stokes modes, the mixing and target mode energies begin to decrease immediately after threshold with no plateau. Now the only important terms in the mixing mode amplitude are the external beam and the molecular modulation term, as originally expected.

Now that the off-resonant SRS has been suppressed, all four modes have nearly-constant complex amplitude at steady state. If we approximate them as constant, we can solve for the amplitudes of the target and mixing modes in terms of the pump and Stokes mode amplitudes. Using the graphical algorithm for determining relevant terms described above, we can solve for...
Fig. 6. Mode energies (left) and mixing mode amplitude composition (right) when the Raman linewidth has been artificially reduced by a factor of $10^3$. The mixing and target mode energies now drop off without a plateau. There are only two important contributions to the mixing mode amplitude: the launched mixing beam and the molecular modulation term $TPS \rightarrow M$ (which is now purely real). These two terms together entirely compose the mixing mode amplitude at steady-state (the two green curves overlap). The reduction of the Raman linewidth has removed the off-resonant SRS between the mixing mode and other modes that was present in Fig. 5.

the target mode amplitude:

$$\hat{E}_T = 0 = - D_T E_T + i G_{MPT} E_M E_S E_P$$

(10)

$$E_T = i \frac{G_{MPT}}{D_T} E_M E_S E_P.$$  

(11)

We can use Eq. (11) to solve for $E_M$, arriving at a form that only depends on $E_P$ and $E_S$:

$$\hat{E}_M = 0 = - D_M E_M + P_M \sqrt{S_M} + i G_{TPSM} E_T E_P E_S$$

(12)

$$0 = - D_M E_M + P_M \sqrt{S_M} + i^2 G_{TPSM} \frac{G_{MPT}}{D_T} E_M E_S E_P E_P E_S$$

(13)

$$E_M = \frac{P_M \sqrt{S_M}}{D_M + G_{TPSM} G_{MPT} |E_P|^2 |E_S|^2 / D_T}.$$  

(14)

When $|E_S|^2$, which is proportional to the Stokes mode energy, becomes large, the second term in the denominator dominates the first and $E_M$ begins to decrease. This is the root cause of the decrease in modulation efficiency: the mixing mode amplitude decreases like $|E_S|^2$, which causes the target mode amplitude to go like $E_T \sim E_M E_S^* \sim E_S^* / |E_S|^2 \sim 1/E_S$ (from Eq. (11)).

This behavior is similar to pump mode clamping in cascaded SRS, where the behavior of a mode that is not being externally pumped causes a change in the form of the evolution equation for another mode that changes its qualitative dynamics. The off-resonant SRS terms, despite their apparently-detritual behavior in Fig. 5, provide extra paths for energy flow that limit the effect of the clamping-like behavior described by Eq. (14) until the Stokes mode reaches much higher energy. This causes the plateau that we observe in the modulation efficiency in Fig. 3. An
observer in the lab would see the launched mixing beam have higher transmission through the fiber as the launched pump power increases, much like the launched pump beam itself begins to have higher transmission due to traditional SRS clamping.

Although this clamping-like effect prevents the modulation efficiency from scaling like the Stokes amplitude as we initially expected, the modulation efficiencies in the plateau region are still very high. Even the simple single-fiber coupling setup gives an efficiency of approximately 0.1% at $\lambda = 800\,\text{nm}$, several orders of magnitude higher than our previous modulator. Critical coupling could boost the efficiency to nearly 10%. The existence of the plateau means that the modulation efficiency should be quite stable to fluctuations in the launched pump power. The MMM will also never need much more launched pump power than the threshold power.

### 3.2. Dependence on four-mode detuning

Bringing the four-mode detuning $\Delta_{MSPT}$ down to less than a mode linewidth $\Gamma_q = \omega_q/Q_q$ (a few MHz for optical frequencies if $Q_q \approx 10^8$) is necessary for high-efficiency operation. Figure 7 shows what happens when $\Delta_{MSPT}$ (which is equal to the target mode detuning $\delta_T$ in that figure) becomes larger than the target mode linewidth $\Gamma_T$. As $\Delta_{MSPT}$ grows, the exponential phase in the relevant terms in Eq. (7) becomes too fast to effectively drive the target mode, and the target mode energy (and therefore output power) begins to decrease like $1/(\Delta_{MSPT}/\Gamma_T)^2$. Equivalently, when this process fails to conserve energy (i.e., when $\hbar\Delta_{MSPT}$ is non-zero), the steady-state efficiency decreases. However, as long as $\Delta_{MSPT} < \Gamma_T$, there is no significant change in $\epsilon$ from perfect four-mode resonance.

![Fig. 7. The modulation efficiency as the target mode frequency $\omega_T$ is detuned from perfect four-mode resonance. Since the other mode frequencies are constant, $\delta_T = \Delta_{MSPT}$, which appears in the molecular modulation terms in Eq. (7). As the detuning becomes larger than the target mode linewidth $\Gamma_T$ (a few MHz for each $\lambda_M$), the modulation efficiency begins to decrease.](image)

Because the mode structure in a microresonator is generally quite irregular, it is important to consider how we could achieve low four-mode detuning in an actual device. One possible method is that the resonator’s mode frequencies could be tuned via the material’s thermo-optic response (the change in index of refraction $n(\omega, T)$ with temperature $T$). The first-order thermo-optic coefficient in silica is $\alpha_n = 1.09 \times 10^{-5}\,\text{K}^{-1}$ [51]. For small changes in the index of refraction, the resonator’s mode frequencies will change by the same fraction that the index of refraction at...
that frequency changes [36]:

\[
\frac{\partial \omega}{\partial T} \approx \frac{\omega}{n(\omega, T)} \frac{\partial n(\omega, T)}{\partial T} = \frac{\omega}{n(\omega, T)} \alpha_n.
\]

(15)

Thus we have

\[
\frac{\partial \Delta_{MSPT}}{\partial T} \approx \alpha_n \left( \frac{\omega_M}{n(\omega_M, T)} - \frac{\omega_S}{n(\omega_S, T)} + \frac{\omega_P}{n(\omega_P, T)} - \frac{\omega_T}{n(\omega_T, T)} \right) .
\]

(16)

If we use \(\lambda_P = 1064\text{nm}\) and \(\lambda_M = 800\text{nm}\), assume that \(\omega_T = \omega_M + 2\pi \times 12\text{THz}\) and \(\omega_S = \omega_P - 2\pi \times 12\text{THz}\), and use the Sellmeier coefficients for silica at 20 °C [52], we find that \(\frac{\partial \Delta_{MSPT}}{\partial T} \approx 2.0\text{MHz K}^{-1}\). With \(\lambda_M = 532\text{nm}\), \(\frac{\partial \Delta_{MSPT}}{\partial T} = 9.1\text{MHz K}^{-1}\).

This calculation indicates that the four-mode detuning could be adjusted by at least a few tens of MHz by a standard thermoelectric controller for any mixing wavelength in the optical region of the spectrum. Although the free spectral range in microresonators is generally at least a few GHz, the actual mode spacing (i.e., the distance to the mode with the closest frequency, not the next mode in the same family) in a non-ideal device can be much denser. For example, by giving a spherical microresonator a small eccentricity to lift the degeneracy of the whispering gallery modes, the actual spacing between adjacent modes could be reduced to a few tens of MHz [53], meaning the four-mode detuning should never be off by more than a few tens of MHz to begin with. However, one may still need to try many different four-mode sets to find a set with high modulation efficiency (due to the mode-dependent coupling factors).

3.3. Dependence on mixing wavelength

One of the key advantages of the MMM is that, depending on the size, geometry, and material of the microresonator, modes could be available over a wide range of wavelengths. Figure 8 shows how the modulation efficiency varies with the wavelength of the mixing mode. From the left panel we see that in the pump-critical case the modulation efficiency decreases dramatically

**Fig. 8.** The modulation efficiency \(\epsilon = P_T/S_M\) as a function of the launched pump power \(S_P\) and the mixing mode wavelength \(\lambda_M\), with the pump mode critically coupled (left) or all modes critically coupled (right). The dominant factor in the decreasing modulation efficiency on the left is the increasingly-non-critical coupling between the fiber and the mixing mode as \(\lambda_M\) is moved further from the pump wavelength \(\lambda_P = 1064\text{nm}\). On the right, where all modes are critically coupled, \(\epsilon\) is essentially independent of \(\lambda_M\).
as the separation between the pump and mixing wavelengths increases because the mixing and target modes become poorly coupled to the fiber (i.e., \( Q_M \) is not close to \( Q_C \)). However, if we can critically couple all of the modes, \( \epsilon \) becomes largely independent of \( \lambda_M \) throughout the optical spectrum, as illustrated in the right panel.

In reality, we would still expect some variation with \( \lambda_M \) because the mixing and target mode volumes and four-mode overlaps will vary, as discussed earlier. Nevertheless, as long as the MMM is built with the capability to adjust the coupling scheme and resonator temperature appropriately, the same device should be able to modulate efficiently over a large, continuous range of mixing beam wavelengths.

### 3.4. Dependence on intrinsic quality factor

Another factor that we could control is the intrinsic quality factor of the microsphere. So far, we have assumed that it is fixed at \( Q^I = 10^8 \). However, microspheres could be produced with either smaller or larger intrinsic quality factors than that. The results of simulations where we vary the intrinsic quality factor are shown in Fig. 9.

![Fig. 9.](image)

| Intrinsic Quality Factor \( Q^I \) | Launched Pump Power \( S_P \) (mW) |
|----------------------------------|-----------------------------|
| \( 10^{-3} \)                    | \( 10^{-1} \)                |
| \( 10^{-2} \)                    | \( 10^{-3} \)                |
| \( 10^{-1} \)                    | \( 10^{-5} \)                |
| \( 10^{1} \)                    | \( 10^{-7} \)                |
| \( 10^{2} \)                    | \( 10^{-9} \)                |

When the intrinsic quality factor is increased at fixed launched pump power the modulation efficiency decreases. Increasing the intrinsic quality factor is essentially the same as shifting the curves in Fig. 3 to lower pump power: threshold decreases, but since we are at fixed launched pump power, we are now further out on the curve, where we already know the modulation efficiency begins to decrease.

### 3.5. Dependence on fiber separation

So far, we have been working with two coupling schemes: either there is a single fiber critically coupled to the pump mode, or all of the modes are critically coupled. In this section, we will use a single fiber as in the first case, but we will change its distance from the microsphere (the fiber separation \( s \)). As \( s \) decreases, the coupling quality factor \( Q^C \) decreases at all wavelengths, meaning that light can more easily move between the fiber and the microsphere. This scheme is the easiest to achieve experimentally, requiring no special modifications of the fiber or microsphere.

The results, shown in Fig. 10, show that the modulation efficiency tends to increase as \( Q^C \) decreases, with no special behavior near the critical separations. This is a result of the target mode not being directly pumped: we do not need to balance input and output power to get critical
coupling, we just need to maximize the output power. Decreasing $Q_C$ lets more light leave the target mode and go to the fiber. Simultaneously, the threshold pump power is increasing, but this is not necessarily detrimental, because we know from Sec. 3.1 that there is no reason to go far beyond threshold. So we have an indirect route to increasing the modulation efficiency by increasing the launched pump power (which eluded us in Sec. 3.1): if we have spare pump power, we can move the fiber closer to the microsphere to get higher efficiency.

Fig. 10. The modulation efficiency $\epsilon = P_T/S_M$ as a function of the launched pump power $S_P$ and the fiber separation $s$. Horizontal dashed lines are the fiber separations where each mode is critically coupled (but note that there is no special behavior at these separations). The pump and mixing wavelengths are $\lambda_P = 1064$nm and $\lambda_M = 800$nm, and the fiber taper radius is 1 $\mu$m. The coupling quality factors $Q_C$ decrease for all modes as the fiber separation decreases.

These results also indicate that the all-critical coupling scheme is not the coupling scheme that maximizes the modulation efficiency. Determining the ideal scheme will depend on a variety of other design parameters and will involve a detailed search through the coupling and intrinsic quality factor parameter space for each mode (in addition to determining how to achieve those quality factors in practice). We plan to explore this in future work.

4. Conclusions

Optical modulators with a modulation frequency of $\sim 10$ THz and an efficiency exceeding 1% will have important applications in a number of research areas. For example, an MMM could be used to produce coherent light at currently-inaccessible parts of the spectrum, such as in the THz regime. An infrared mixing beam could be frequency-downshifted to efficiently produce THz radiation [54]. Another application is to optical waveform synthesis. Because the modulation efficiency is very large, higher-order Stokes and anti-Stokes sidebands could be produced by sending a mixing beam through several MMMs in series. These sidebands would form a broad spectrum, which could then be used to synthesize optical waveforms at femtosecond time scales.

Microsphere-based molecular modulators could represent a major step toward realizing these applications because they can potentially achieve extremely high modulation efficiency over a wide range of mixing beam wavelengths. Compared to our previous modulator, they also require less pump power, are more stable, are easier to operate, and are readily interoperable with other fiber-coupled devices.

There are still many challenges to overcome to build a functioning MMM: determining the optimal geometry and material for the microresonator, finding the best coupling scheme,
characterizing the time-dependent behavior during device startup, and understanding how the presence of additional excited modes and even intermode coupling will impact the modulation efficiency [38], to name just a few. Nevertheless, MMMs are a significantly better platform to work with than our previous modulator design, with a much higher ceiling for performance and a much lower floor for technical problems.

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