Maximal spontaneous photon emission and energy loss from free electrons

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Free-electron radiation such as Cerenkov4, Smith–Purcell2 and transition radiation24,25 can be greatly affected by structured optical environments, as has been demonstrated in a variety of polaritonic5,6, photonic-crystal1 and metamaterial8–10 systems. However, the amount of radiation that can ultimately be extracted from free electrons near an arbitrary material structure has remained elusive. Here we derive a fundamental upper limit to the spontaneous photon emission and energy loss of free electrons, regardless of geometry, which illuminates the effects of material properties and electron velocities. We obtain experimental evidence for our theory with quantitative measurements of Smith–Purcell radiation. Our framework allows us to make two predictions. One is a new regime of radiation operation—at subwavelength separations, slower (non-relativistic) electrons can achieve stronger radiation than fast (relativistic) electrons. The other is a divergence of the emission probability in the limit of lossless materials. We further reveal that such divergences can be approached by coupling free electrons to photonic bound states in the continuum13–15. Our findings suggest that compact and efficient free-electron radiation sources from microwaves to the soft X-ray regime may be achievable without requiring ultrahigh accelerating voltages.

The Smith–Purcell effect epitomizes the potential of free-electron radiation. Consider an electron at velocity β = v/c traversing a structure with periodicity a; it generates far-field radiation at wavelength λ and polar angle θ, dictated by2:

\[ \lambda = \frac{a}{m} \left( \frac{1}{\beta} \cos \theta \right) \]  

where m is the integer diffraction order. The absence of a minimum velocity in equation (1) offers prospects for threshold-free and spectrally tunable light sources, spanning from microwave and terahertz14–16, across visible17–19, and towards X-ray20 frequencies. In stark contrast to the simple momentum-conservation determination of wavelength and angle, there is no unified yet simple analytical equation for the radiation intensity. Previous theories offer explicit solutions only under strong assumptions (for example, assuming perfect conductors or employing effective medium descriptions) or for simple, symmetric geometries21–23. Consequently, heavily numerical strategies are often an unavoidable resort14,24. In general, the inherent complexity of the interactions between electrons and photonic media have prevented a more general understanding of how pronounced spontaneous electron radiation can ultimately be for arbitrary structures, and consequently, how to design the maximum enhancement for free-electron light-emitting devices.

We begin our analysis by considering an electron (charge −e) of constant velocity v traversing a generic scatterer (plasmonic or dielectric, finite or extended) of arbitrary size and material composition, as in Fig. 1a. The free current density of the electron, \( J(r, t) = -\delta(x - vt) \), generates a frequency-dependent (c−ω convention) incident field:

\[
F_{\text{inc}}(r, \omega) = \frac{eK_e}{2\pi ev_0} \left[ \delta(x)K_0(k_\rho) - \hat{p}_x K_1(k_\rho) \right] 
\]

written in cylindrical coordinates (x, ρ, φ); here, \( K_n \) is the modified Bessel function of the second kind, \( k_\rho = \omega/\gamma \) and \( k_v = \sqrt{k_v^2 - \omega^2} = k|\beta|/\gamma \) (k = αc, free-space wavevector; \( \gamma = 1/\sqrt{1 - \beta^2} \), Lorentz factor). Hence, the photon emission and energy loss of free electrons can be treated as a scattering problem: the electromagnetic fields \( F_{\text{inc}} = (E_{\text{inc}}, Z_0 H_{\text{inc}})^T \) (for free-space impedance \( Z_0 \)) are incident on a photonic medium with material susceptibility \( \chi \) (a 6×6 tensor for a general medium), causing both absorption and far-field scattering—that is, photon emission—that together comprise electron energy loss (Fig. 1a).

As recently shown in refs 27–29, for a generic electromagnetic scattering problem, passivity—the condition that polarization currents do no work—constrains the maximum optical response from a given incident field. Consider three power quantities derived from \( F_{\text{inc}} \) and the total field \( F \) within the scatterer volume \( V \): the total power lost by the electron, \( P_{\text{loss}} = -(1/2)\text{Re} \int_V J \cdot \mathcal{E} dV = (e\omega/2)\text{Im} \int_V F_{\text{inc}}^\dagger \mathcal{F} FdV \), the power absorbed by the medium, \( P_{\text{abs}} = (e\omega/2)\text{Im} \int_V F^\dagger \mathcal{F} FdV \), and their difference, the power radiated to the far field, \( P_{\text{rad}} = P_{\text{loss}} - P_{\text{abs}} \). Treating \( F \) as an independent variable, the total loss \( P_{\text{loss}} \) is a linear function of \( F \), whereas the fraction that is dissipated is a quadratic function of \( F \). Passivity requires non-negative radiative power, represented by the inequality \( P_{\text{abs}} < P_{\text{loss}} \) which in this framework is therefore a convex constraint on any response function. Constrained maximization (see Supplementary Section 1) of the energy-loss and photon-emission power quantities, \( P_{\text{loss}} \) and \( P_{\text{rad}} \), directly yields the limits:

\[
P_{\text{f}}(\omega) \leq \frac{E_0\omega^2}{2} \int_V F_{\text{inc}}^\dagger (\text{Im} \mathcal{F})^{-1} F_{\text{inc}} dV 
\]
Absorption

\( \kappa \) reflects the influence of the material choice, which constrains the maximum material response of the impact parameter limit (see Supplementary Section 2) that highlights the pivotal role of isolation vs. metallic media. The simplified, leading to a general closed-form solution based on the electron velocity \( \beta \) is the fine-structure constant. Equation (4) imposes, without solving Maxwell’s equations, a maximum rate of photon generation based on the electron velocity \( \beta \) (through \( k_e \) and \( \kappa_e \)), the material composition \( \chi(\tau) \) and the volume \( V \).

The limit in equation (4) can be further simplified by removing the shape dependence of \( V \), since the integrand is positive and is thus bounded above by the same integral for any enclosing structure. A scatterer separated from the electron by a minimum distance \( d \) can be enclosed within a larger concentric hollow cylinder sector of inner radius \( d \) and outer radius \( \infty \). For such a sector (height \( L \) and opening azimuthal angle \( \varphi \in (0, 2\pi) \)), equation (4) can be further simplified, leading to a general closed-form shape-independent limit (see Supplementary Section 2) that highlights the pivotal role of the impact parameter \( \kappa d \):

\[
\Gamma(\omega) \leq \frac{\alpha_2 e}{2 c \omega} \int_0^\infty \frac{|\chi|^2}{1 + \beta^2} [\kappa_0^2 K_0^2(k_e \rho) + \kappa_e^2 R^2 K_1^2(k_e \rho)] dV
\]

where \( \alpha_2 \) is the fine-structure constant. Equation (4) imposes, without solving Maxwell’s equations, a maximum rate of photon generation based on the electron velocity \( \beta \) (through \( k_e \) and \( \kappa_e \)), the material composition \( \chi(\tau) \) and the volume \( V \).

A surprising feature is the limits of equations (4), (5a) and (5b) is their prediction for optimal electron velocities. As shown in Fig. 1c, when electrons are in the far field of the structure \( \kappa d \gg 1 \), their interaction is characterized by the dominant fields that are the incident field \( E_{\text{inc}} \) and the scattered field \( E_{\text{scat}} \), which are perpendicular to each other. The electric field amplitude of the incident field is given by:

\[
E_{\text{inc}}(r) = E_0 \frac{\sin(k d)}{k d}
\]

where \( E_0 \) is the electric field amplitude of the plane wave and \( k \) is the wave vector of the incident light. The scattered field is given by:

\[
E_{\text{scat}}(r) = \frac{\chi}{k_e^2} \frac{\sin(k d)}{k d} \nabla \cdot \mathbf{E}_{\text{inc}} - \frac{\chi}{k_e^2} \frac{\cos(k d)}{k d} \nabla \times \mathbf{E}_{\text{inc}}
\]

where \( \mathbf{E}_{\text{inc}} \) is the electric field of the incident light and \( \chi \) is the susceptibility of the material. The electric field amplitude of the scattered field is given by:

\[
E_{\text{scat}}(r) = E_0 \frac{\sin(k d)}{k d} \left( \frac{\chi}{k_e^2} \frac{\sin(k d)}{k d} \nabla \cdot \mathbf{E}_{\text{inc}} - \frac{\chi}{k_e^2} \frac{\cos(k d)}{k d} \nabla \times \mathbf{E}_{\text{inc}} \right)
\]

The limits of equations (4), (5a) and (5b) are completely general; they set the maximum photon emission and energy loss of an electron beam coupled to an arbitrary photonic environment in either the non-retarded or retarded regimes, given only the beam properties and material composition. The key factors that determine maximal radiation are identified: intrinsic material loss (represented by \( \eta^2 \)), electron velocity \( \beta \) and impact parameter \( \kappa d \). The metric \( \frac{\beta^2}{2} \eta \) reflects the influence of the material choice, which depends sensitively on the radiation wavelength (Fig. 1b). The electron velocity \( \beta \) also appears implicitly in the impact parameter \( \kappa d \). The impact parameter \( \kappa d \) reflects the influence of the Lorentz contraction \( d/\beta \), a well-known feature of both electron radiation and acceleration.

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\]

where \( \mathbf{E}_{\text{inc}} \) is the electric field of the incident light and \( \chi \) is the susceptibility of the material. The electric field amplitude of the scattered field is given by:

\[
E_{\text{scat}}(r) = E_0 \frac{\sin(k d)}{k d} \left( \frac{\chi}{k_e^2} \frac{\sin(k d)}{k d} \nabla \cdot \mathbf{E}_{\text{inc}} - \frac{\chi}{k_e^2} \frac{\cos(k d)}{k d} \nabla \times \mathbf{E}_{\text{inc}} \right)
\]
Two scenarios are considered: in Fig. 1d, an electron traverses the centre of an annular Au bowtie antenna and undergoes antenna-enabled transition radiation \((\eta \approx 0.07\%\)) while, in Fig. 1e, an electron traverses a Au grating, undergoing Smith–Purcell radiation \((\eta \approx 0.9\%\)). In both cases, the numerical results closely trail the upper limit at the considered wavelengths, showing that the limits can be approached or even attained with modest effort.

Next, we specialize in the canonical Smith–Purcell set-up illustrated in Fig. 1e inset. This set-up warrants a particularly close study, given its prominent historical and practical role in free-electron radiation. Aside from the shape-independent limit (equations (5a) and (5b)), we can find a sharper limit (in per unit length for periodic structure) specifically for Smith–Purcell radiation using rectangular gratings of filling factor \(A\) (see Supplementary Section 3).

\[
\frac{dG(\omega)}{dx} \leq \frac{a^2}{2\pi} |k|^2 \Lambda G(\beta, kd)
\]

The function \(G(\beta, kd)\) is an azimuthal integral (see Supplementary Section 3) over the Meijer G-function \(G_{1,3}\) (ref. \(^{20}\)) that arises in the radial integration of the modified Bessel functions \(K_\nu\). We emphasize that equation (6) is a specific case of equation (4) for grating structures without any approximations and thus can be readily generalized to multi-material scenarios (see Supplementary equation (37)).

The grating limit (equation (6)) exhibits the same asymptotics as equations (5a) and (5b), thereby reinforcing the optimal-velocity predictions of Fig. 1c. The \((\beta, kd)\) dependence of \(G\) (see Fig. 2a) shows that slow (fast) electrons maximize Smith–Purcell radiation in the small (large) separation regime. We verify the limit predictions by comparison with numerical simulations: at small separations (Fig. 2b), radiation and energy loss peak at velocity \(\beta \approx 0.15\), consistent with the limit maximum; at large separations (Fig. 2c), both the limit and the numerical results grow monotonically with \(\beta\).

The derived upper limit also applies to Cherenkov and transition radiation, as well as bulk loss in electron energy-loss spectroscopy. For these scenarios where electrons enter material bulk, a subtlety arises for the field divergence along the electron’s trajectory (\(\rho = 0\) in equation (2)) within a potentially lossy medium. This divergence, however, can be regularized by introducing natural, system-specific momentum cutoffs\(^{27}\), which then directly permits the application of our theory (see Supplementary Section 6). Meanwhile, there exist additional competing interaction processes (for example, electrons colliding with individual atoms). However, they typically occur at much smaller length scales.

We perform quantitative experimental measurement of Smith–Purcell radiation to directly probe the upper limit. Figure 3a shows our experimental set-up (see Methods and Supplementary Section 7 for details). A one-dimensional (1D) 50%-filling-factor grating (Au-covered single-crystalline Si)—the quintessential Smith–Purcell set-up—is chosen as a sample, and shown by scanning electron microscope (SEM) images in Fig. 3b,c. Free electrons pass above and impinge onto the sample at a grazing angle of 1.5° under 10 to 20 kV acceleration voltages.

Figure 3d depicts our measurements of first-order \(m = 1\) Smith–Purcell radiation appearing at wavelengths between 500 and 750 nm. In quantitative agreement with equation (1) evaluated at the normal emission angle (solid lines), the measured radiation spectra (dots) blueshift with increasing electron velocity. Notably, we experimentally obtain the absolute intensity of the collected radiation via a calibration measurement (see Supplementary Section 7). The upper limits (equation (4)) for the surface-normal emission wavelengths \((\lambda = a/\beta)\) are evaluated at the centre of the interaction region (height \(\approx 140\) nm \((kd \approx 1.5)\), varying with beam energy), and is shown with shading in Fig. 3d to account for the thickness uncertainty \(\pm 1.5\) nm. The envelope spanned by the measurement peaks follows the upper-limit lineshape across the visible spectrum; both the theoretical limit and the measured intensities peak near 550 nm and decrease in a commensurate manner for other wavelengths. This lineshape originates from two competing factors. At shorter wavelengths, the material factor \(|k|^2/1m\gamma\) decreases significantly for both Au and Si (Fig. 1c), which accounts for the reduced radiation intensity. At longer wavelengths, the major constraint becomes the less efficient interaction between the electrons and the structure, as the electron-beam diameters increase for the reduced brightness of the electron gun (tungsten) at lower acceleration voltages (see Supplementary Section 7). These pieces of experimental evidence for the upper limit are at \(kd \approx 1.5\) (estimated from a geometrical ray-tracing model; see Supplementary Section 7), where fast electrons are still preferred (Fig. 2a). To further confirm our theory, we also conduct a near-infrared Smith–Purcell experiment (Supplementary Section 8) at \(kd \approx 1\), where the envelope lineshape of the emission spectra again follows our prediction. We also obtain complementary supporting evidence (extracted from a recent work\(^{23}\)) for our slow-electron-efficient prediction (see Supplementary Section 9).

\(\begin{align*}
\text{Fig. 2} & \text{ Optimal electron velocities for maximal Smith–Purcell radiation.} \\
\text{a. Behaviour of } G(\beta, kd), \text{ equation (6)}, \text{ whose maxima indicate separation-dependent optimal electron velocities. Here } G \text{ is normalized between 0 and 1 for each separation. The limit yields sharply contrasting predictions: slow electrons are optimal in the near field (kd } \ll 1 \text{) and fast electrons are optimal in the far field (kd } \gg 1 \text{). b, c. Energy loss (red) and radiation (blue) rates (circles: full-wave simulations; lines: grating limit, equation (6); shading: shape-independent limit, equations (5a) and (5b)) at two representative near/far-field separation distances (white dashed slices in a).}
\end{align*}\)
Finally, we turn our attention to an ostensible peculiarity of the limits: equation (4) evidently diverges for lossless materials \((\text{Im} \chi \rightarrow 0)\), seemingly providing little insight. On the contrary, this divergence suggests the existence of a mechanism capable of strongly enhancing Smith–Purcell radiation. Indeed, by exploiting high-Q resonances near bound states in the continuum (BICs)\(^{13}\) in photonic crystal slabs, we find that Smith–Purcell radiation can be enhanced by orders of magnitude, when specific frequency-, phase- and polarization-matching conditions are met.

A 1D silicon \((\chi = 11.25)\)-on-insulator \((\text{SiO}_2, \chi = 1.07)\) grating interacting with a sheet electron beam illustrates the core conceptual idea most clearly. The transverse electric (TE) \((E_x, H_y, E_z)\) band structure (lowest two bands labelled \(T_E\) and \(T_E\)), matched polarization for a sheet electron beam (supplementary equation S41b)), is depicted in Fig. 4b along the \(\Gamma-X\) direction. Folded electron wavevectors, \(k = \omega / v\), are overlaid for two distinct velocities (blue and green). Strong electron–photon interactions are possible when the electron and photon dispersions intersect: for instance, \(k_x\) and the \(T_E\) band intersect (grey circles) below the air light cone (light yellow shading). However, these intersections are largely impractical: the \(T_E\) band is evanescent in the air region, precluding free-space radiation. Still, analogous ideas, employing similar partially guided modes, such as spoof plasmons\(^{33}\), have been explored for generating electron-enabled guided waves\(^{34,35}\).

To overcome this deficiency, we theoretically propose a new mechanism for enhanced Smith–Purcell radiation: coupling of electrons with BICs\(^{13}\). The latter have the extreme quality factors of guided modes but are, crucially, embedded in the radiation continuum, guaranteeing any resulting Smith–Purcell radiation into the far field. By choosing appropriate velocities \(\beta = a / m \lambda\) (\(m\) being any integer; \(\lambda\) being the BIC wavelength) such that the electron line (blue or green) intersects the \(T_E\) mode at the BIC (red square in Fig. 4b), the strong enhancements of a guided mode can be achieved in tandem with the radiative coupling of a continuum resonance. In Fig. 4c, the incident fields of electrons and the field profile of the BIC are shown complete confinement without radiation, unlike conventional multipolar radiation modes (see Supplementary Fig. 9). The Q values of the resonances are also provided near a symmetry-protected BIC\(^{13}\) at the \(\Gamma\) point. Figure 4d,e demonstrates the velocity tunability of BIC-enhanced radiation—as the phase matching approaches the BIC, a divergent radiation rate is achieved.

The BIC-enhancement mechanism is entirely accordant with our upper limits. Practically, silicon has non-zero loss across the visible and near-infrared wavelengths. For example, for a period of \(a = 676\) nm, the optimally enhanced radiation wavelength is \(\approx 1,050\) nm, at which \(\chi_x \approx 11.25 + 0.001\) (ref.\(^{36}\)). For an electron–structure separation of 300 nm, we theoretically show in Fig. 4f the strong radiation enhancements (>3 orders of magnitude) attainable by BIC-enhanced coupling. The upper limit (shaded region; 2D analogue of equation (4); see Supplementary Section 10) attains extremely large values due to the minute material loss \((|\chi|^2 / \text{Im} \chi \approx 10^3)\); nevertheless, BIC-enhanced coupling enables the radiation intensity to closely approach this limit at several resonant velocities. In the presence of an absorptive channel, the maximum enhancement occurs at a small offset from the BIC where the envelope (peak outline) of the measured spectra (dots) follows the theoretical upper limit (shaded to account for fabrication tolerance; calculated at each wavelength with the corresponding electron velocity for surface-normal radiation).

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**Fig. 3 | Experimental probing of the upper limit.** a, Experimental set-up. OBJ, objective (numerical aperture of 0.3); BS, beamsplitter; SP, spectrometer; CAM, camera. b, c, SEM images of the structure in top view (b) and cross-sectional view (c). d, Quantitative measurement of Smith–Purcell radiation (inset: camera image of the radiation). The solid lines mark the theoretical radiation wavelengths at the normal angle (equation (1)). The envelope (peak outline) of the measured spectra (dots) follows the theoretical upper limit (shaded to account for fabrication tolerance; calculated at each wavelength with the corresponding electron velocity for surface-normal radiation).
In closing, we have theoretically derived and experimentally probed a universal upper limit to the energy loss and photon emission from free electrons. The limit depends crucially on the impact parameter $x, d$, but not on any other detail of the geometry. Hence, our limit applies even to the most complex metamaterials and metasurfaces, given only their constituents. Surprisingly, in the near field, slow electrons promise stronger radiation than relativistic ones. The limit predicts a divergent radiation rate as the material loss rate goes to zero, and we show that BIC resonances enable such staggering enhancements. This is relevant for the generation of coherent Smith–Purcell radiation14,34,35. The long lifetime, spectral selectivity and large field enhancement near a BIC can strongly bunch electrons, allowing them to radiate coherently at the same desired frequency, potentially enabling low-threshold Smith–Purcell free-electron lasers. The combination of this mechanism and the optimal velocity prediction reveals prospects of low-voltage yet high-power free-electron radiation sources. In addition, our findings demonstrate a simple guiding principle to maximize the signal-to-noise ratio for electron energy-loss spectroscopy through an optimal choice of electron velocity, enabling improved spectral resolution.

The upper limit demonstrated here is in the spontaneous emission regime for constant-velocity electrons, and can be extended to the stimulated regime by suitable reformulation. Stronger electron–photon interactions can change electron velocity by a non-negligible amount that alters the radiation. If necessary, this correction can be perturbatively incorporated. In the case of external optical pumping37, the upper limit can be revised by redefining the incident field as the summation of the electron incident field and the external optical field. From a quantum mechanical perspective, this treatment corresponds to stimulated emission from free electrons, which multiplies the limit by the number of photons in that radiation mode. This treatment could also potentially translate our limit into a fundamental limit for particle acceleration38,39, which is the time-reversal of free-electron energy loss and which typically incorporates intense laser pumping. In the multi-electron scenario, the radiation upper limit will be obtained in the case of perfect bunching, where all electrons radiate in phase. In this case, our single-electron limit should be multiplied by the number of electrons to correct for the superradiant nature of such coherent radiation.

Methods

Methods, including statements of data availability and any associated accession codes and references, are available at https://doi.org/10.1038/s41567-018-0180-2

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Author contributions

Y.Y., O.D.M., I.K. and M.S. conceived the project. Y.Y. developed the analytical models and numerical calculations. A.M. prepared the sample under study. Y.Y., A.M., C.R.-C., S.E.K. and I.K. performed the experiment. Y.Y., T.C. and O.D.M. analysed the agreement no. 328853CMC-BSiCS. Programme of the European Research Council (FP7– Marie Curie IOP) under grant agreement no. 328853MC–B5SIC.

Competing interests

The authors declare no competing interests.

Additional information

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Methods

Fourier transform convention. Throughout the paper, we adopt the following Fourier transform conventions

\[
f(\omega) \equiv \int f(t) e^{i\omega t} dt, \quad f(t) \equiv \frac{1}{2\pi} \int f(\omega) e^{-i\omega t} d\omega
\]

(7)

\[
g(k) \equiv \int f(r) e^{-ikr} dr, \quad g(r) \equiv \frac{1}{(2\pi)^3} \int g(k) e^{i\mathbf{k} \cdot \mathbf{r}} d\mathbf{k}
\]

(8)

Numerical methods. The photonic band structure in Fig. 4b is calculated via the eigenfrequency calculation in COMSOL Multiphysics. Numerical radiation intensities (Figs. 1d,e, 2b,c, 3d and 4d–f) are obtained via the frequency-domain calculation in the radiofrequency module in COMSOL Multiphysics. A surface (for 3D problems) or line (for 2D problems) integral on the Poynting vector is calculated to extract the radiation intensity at each frequency.

Experimental set-up and sample fabrication. Our experimental set-up comprises a conventional SEM with the sample mounted perpendicular to the stage. A microscope objective was placed on the SEM stage to collect and image the light emission from the surface. The collected light was then sent through a series of free-space optical elements, enabling simultaneous measurement of the spectrum and of the spatial radiation pattern.

The SEM used for the experiment was a JEOL JSM-6010LA. Its energy spread at the gun exit was in the range 1.5 to 2.5 eV for the range of acceleration voltages considered in this paper. The SEM was operated in spot mode, which we controlled precisely to align the beam so that it passes tangentially to the surface near the desired area of the sample. A Nikon TU Plan Fluor 10x objective was used to collect light from the area of interest. The monochrome image of the radiation was taken using a Hamamatsu CCD (charge-coupled device). The spectrometer used was an Action SP-2360-2300i with a low-noise Princeton Instruments Pixis 400 CCD.

A 1D grating (Au-covered single-crystalline Si: periodicity, 140 nm; filling factor, 50%; patterned Si thickness, 53 ± 1.5 nm; Au thickness 44 ± 1.5 nm) was used as the sample in our experiment. The original nanopatterned linear silicon stamp was obtained from LightSmyth Technologies and coated using an electron beam evaporator with a 2 nm Ti adhesion layer and 40 nm of Au at 10⁻⁷ torr. The sample was mounted inside the SEM chamber to enable the alignment of free electrons to pass in close proximity to the stamps. The emitted light was coupled out of the SEM chamber to a spectrometer, while a camera was used to image the surface of the sample.

Data availability. The data that support the plots within this paper and other findings of this study are available from the corresponding author upon reasonable request.
Maximal spontaneous photon emission and energy loss from free electrons

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SUPPLEMENTARY INFORMATION

Maximal Spontaneous Photon Emission and Energy Loss from Free Electrons

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1. **GENERAL OPTICAL RESPONSE LIMIT FRAMEWORK**

Intuitively, to impose the general limits on the energy loss and photon emission (cathodoluminescence) from free electrons, we consider an arbitrary scatterer embedded in a possibly heterogeneous background. Passivity, which implies the absence of gain and that polarization currents do no work \([1]\) requires that the absorbed \((P_{\text{abs}})\) and scattered \((P_{\text{rad}})\) powers by the target body are non-negative. On the other hand, their sum, the electron energy loss \((P_{\text{loss}} = P_{\text{abs}} + P_{\text{rad}})\), is given by the real part of the overlap between the electron velocity and the induced field \([2]\), similar to the optical theorem \([3]\).

More broadly, for an arbitrarily shaped 3D scatterer [volume \(V\) and susceptibility tensor \(\chi(r, \omega)\)] impinged by the external incident field \(\mathbf{F}_{\text{inc}} = (\mathbf{E}_{\text{inc}}, Z_0 \mathbf{H}_{\text{inc}})^T\) [for the case of free electrons, see Eq. (2)], the absorption (dissipation) within such a medium is the work done by the total fields \(\mathbf{F}\) on the induced currents, given by the expression

\[
P_{\text{abs}} = \frac{\varepsilon_0\omega}{2} \text{Im} \int_V \mathbf{F}^\dagger \chi \mathbf{F} \, dV. \tag{S1}
\]

On the other hand, the total electron energy loss represents the work done by the incident fields on the induced currents

\[
P_{\text{loss}} = \frac{\varepsilon_0\omega}{2} \text{Im} \int_V \mathbf{F}^\dagger_{\text{inc}} \chi \mathbf{F} \, dV. \tag{S2}
\]

As can be seen, Electron total energy loss and absorption are linear and quadratic function of the fields, respectively. Yet electron energy loss must be greater than absorption (due to the nonnegative scattering noted above), requiring the linear functional to be greater than the quadratic one, a condition that cannot be satisfied for large enough currents. The inequality \(P_{\text{abs}} \leq P_{\text{loss}}\) thereby provides a convex constraint for the optical excitation of free electrons. Thanks to the convex nature of the constraint \(P_{\text{abs}} \leq P_{\text{loss}}\) and the simple expressions of the absorption and energy loss, the optimal response can be solved analytically using variational derivatives, without the requirement of solving the highly nonconvex Maxwell equations, thereby providing general upper-limit expressions for electron energy loss and photon emission without approximation.

To obtain the extremum induced fields/currents for electron energy loss or photon emission (cathodoluminescence), one can take the derivative \(\frac{\partial P_{\tau}(\omega)}{\partial \mathbf{F}^\dagger} = 0\), where \(\tau \in \{\text{rad, loss}\}\). Using the photon emission as an example, by taking

\[
\frac{\partial P_{\text{rad}}(\omega)}{\partial \mathbf{F}^\dagger} = \frac{\partial (P_{\text{loss}} - P_{\text{abs}})}{\partial \mathbf{F}^\dagger} = 0, \tag{S3}
\]
we have

$$\chi^\dagger F^\text{inc}/2i + (\text{Im}\chi)F = 0,$$  \hspace{1cm} (S4)

where $\text{Im}\chi = (\chi - \chi^\dagger)/2i$ is a Hermitian matrix. From Eq. (S4), one readily finds the optimal total field is given by $F_{\text{rad, opt}} = i/2(\text{Im}\chi)^{-1}\chi^\dagger F^\text{inc}$. The optimal field for maximal electron energy loss can be derived in a similar manner. Combining the optimal fields with Eq. (S1) and Eq. (S2), we obtain the upper limit shown in Eq. (3) in the main text.

2. THREE-DIMENSIONAL SHAPE-INDEPENDENT UPPER LIMIT

We first rewrite the three-dimensional general limit equation

$$\Gamma_\tau(\omega) \leq \frac{e^2 \xi_\tau}{8\hbar\varepsilon_0\omega^2\pi^2} \int |\chi|^2 \text{Im} \chi \left[ \kappa^4_\rho K^2_0(\kappa_\rho \rho) + \kappa^2_\rho k^2_\nu K^2_1(\kappa_\rho \rho) \right] dV.$$  \hspace{1cm} (S5)

We assume the structure is made of a single material

$$\Gamma_\tau(\omega) \leq \frac{e^2 \xi_\tau}{8\hbar\varepsilon_0\omega^2\pi^2} \frac{|\chi|^2}{\text{Im} \chi} \int |\chi|^2 \left[ \kappa^4_\rho K^2_0(\kappa_\rho \rho) + \kappa^2_\rho k^2_\nu K^2_1(\kappa_\rho \rho) \right] dV.$$  \hspace{1cm} (S6)

We now simplify the integral

$$\mathcal{K} = \int |\chi|^2 \left[ \kappa^4_\rho K^2_0(\kappa_\rho \rho) + \kappa^2_\rho k^2_\nu K^2_1(\kappa_\rho \rho) \right] dV.$$  \hspace{1cm} (S7)

For an arbitrarily-shaped structure, whether isolated or extended, one can always find a circular concentric hollow cylinder (height $L$, opening azimuthal angle $\psi \in [0, 2\pi]$, minor radius being the electron structure separation, major radius can be finite or infinite) that encloses it. Therefore, we can evaluate the integral in the cylindrical coordinate

$$\mathcal{K} \leq L\psi \int_0^\infty \rho \left[ \kappa^4_\rho K^2_0(\kappa_\rho \rho) + \kappa^2_\rho k^2_\nu K^2_1(\kappa_\rho \rho) \right] d\rho$$

$$= L\psi \int_{x_0}^\infty x \left[ \kappa^2_\rho K^2_0(x) + k^2_\nu K^2_1(x) \right] dx,$$

$$= \frac{x_0^2}{2} \left\{ \kappa^2_\rho \left[ K^2_0(x_0) - K^2_0(0) \right] + k^2_\nu \left[ K_0(x_0)K_2(x_0) - K^2_1(x_0) \right] \right\},$$

$$= \frac{x_0^2}{2} \left\{ k^2_\nu K_0(x_0) \left[ K_2(x_0) - K_0(x_0) \right] - k^2 \left[ K^2_1(x_0) - K^2_0(x_0) \right] \right\},$$

$$= x_0 k^2_\nu K_0(x_0) K_1(x_0) - x_0^2 k^2 \left[ K^2_1(x_0) - K^2_0(x_0) \right],$$

$$\leq x_0 k^2_\nu K_0(x_0) K_1(x_0),$$  \hspace{1cm} (S8)
where \( x_0 = \kappa_p d \). Eq. (S8) corresponds to Eq. (5a) in the maintext. In the derivation above, we use the following relations [4]

\[
\int x K_n^2(x) dx = \frac{x^2}{2} \left[ K_n^2(x) - K_{n-1}(x) K_{n+1}(x) \right], \tag{S9a}
\]

\( K_{-1}(x) = K_1(x) \), \( K_2(x) - K_0(x) = 2K_1(x)/x \), \( K_1(x) > K_0(x) \). \( \tag{S9d} \)

Figure S1. (a) Longitudinal \( E_x \) and (b) transverse \( E_\rho \) incident field amplitudes generated from free electrons in the azimuthal direction \( \rho \) at different velocities \( \beta \).

In the main text, the shape-independent limit has sharply-contrasting prediction on the intensity of luminescence and energy loss of free electrons when they are in either the far or near field [Eq. (5b)]. Fig. S1 shows that the incident fields already exhibit similar property, which naturally translates into the upper limit via the overlap integral [Eq. (4)].

3. SMITH–PURCELL RADIATION UPPER LIMIT IN THREE DIMENSIONS FOR RECTANGULAR GRATINGS

We choose coordinates such that \((vt, y_0, z_0)\) depicts the trajectory of the charged particle. In the cylindrical coordinate \((\rho, \psi, x)\), the current density can be rewritten as

\[
J(r, t) = \frac{-ev}{2\pi\rho} \delta(x - vt)\delta(\rho)\hat{x}.
\]
where \( \Gamma \) is the gamma function.

Fourier transform on Eq. (S10) yields the current density in the frequency domain

\[
\mathbf{J}(r, \omega) = \frac{-e}{2\pi \rho} e^{ik_x x} \delta(\rho) \hat{x},
\]  
(S11)

whose external electromagnetic field is given by [5]

\[
\mathbf{E}_{\text{inc}}(r, \omega) = \frac{e}{4\omega \varepsilon_0} (k_x^2 \hat{x} + ik_y \nabla) H_0^{(1)}(ik_x \rho) e^{ik_x x},
\]

\[
= \frac{e}{4\omega \varepsilon_0} [(k_x^2 - k_y^2) H_0^{(1)}(ik_x \rho) \hat{x} + ik_y d_y H_0^{(1)}(ik_x \rho) \hat{\rho}] e^{ik_y y},
\]

\[
= \frac{e}{2\pi \omega \varepsilon_0} [ik_y^2 K_0(ik_x \rho) \hat{x} - k_x k_y K_1(ik_x \rho) \hat{\rho}] e^{ik_y y}.
\]  
(S12)

where \( H_0^{(1)} \) is the Hankel function of the first kind with zero order. Here we utilize the relation

\[ K_0(z) = \frac{i\pi}{2} H_0^{(1)}(iz), \]

where \( z \) is a real argument. Insert Eq. (S12) into Eq. (3) yields the general three-dimensional limit shown in Eq. (4).

Next we consider Smith–Purcell radiation from rectangular gratings in three dimensions. The volume integral of the evanescent field is given by

\[
\int_V |\mathbf{E}(r)|^2 dV = \frac{e^2}{4\omega^2 \varepsilon_0 \pi^2} \int dx \int_{\pi/2}^{\pi/2} d\psi \int_{\psi/\cos \psi}^{\infty} \rho \, d\rho \, K_0^2(\rho) + \rho \, \rho \, K_1^2(\rho),
\]  
(S13)

Closed-form integral can be obtained by using the relation

\[
\int_{d/\cos \psi}^{\infty} \rho \, d\rho K_0^2(\rho) = \frac{\sqrt{\pi}}{4\rho} G_{1,3}^{3,0} \left( \rho^2 d^2 \sec^2 \psi_{1,0,1}^{3/2} \right),
\]

and

\[
\int_{d/\cos \psi}^{\infty} \rho \, d\rho K_1^2(\rho) = \frac{\sqrt{\pi}}{4\rho} G_{1,3}^{3,0} \left( \rho^2 d^2 \sec^2 \psi_{0,0,2}^{3/2} \right).
\]  
(S15)

Here \( G \) is the Meijer G-function [4, 6] defined as a line integral in the complex plane

\[
G_{\rho,q}^{m,n}(z \mid a_1, \ldots, a_p \mid b_1, \ldots, b_q) = \frac{1}{2\pi i} \int_L \prod_{j=1}^{m} \Gamma(b_j - s) \prod_{j=1}^{n} \Gamma(1 - a_j + s) \prod_{j=m+1}^{q} \Gamma(1 - b_j + s) \prod_{j=n+1}^{p} \Gamma(a_j - s) \, z^s \, ds,
\]  
(S16)

where \( \Gamma \) is the gamma function.

Plug Eq. (S14) and Eq. (S15) into Eq. (4) yields Eq. (6) in the main text

\[
\frac{d\Gamma(\omega)}{dx} \leq \frac{\alpha x}{2\pi c \, \text{Im} \chi} \mathcal{G}(\beta, \kappa) \left. \Lambda \mathcal{G}(\beta, \kappa) \right|^{1/2},
\]  
(S17a)

where

\[
\mathcal{G}(\beta, \kappa) = \mathcal{G}_s(\beta, k_d) + \mathcal{G}_p(\beta, k_d),
\]  
(S17b)

\[
\mathcal{G}_s(\beta, k_d) = \frac{\sqrt{\pi}}{4} \int_{-\pi/2}^{\pi/2} \frac{d\psi}{\beta^2 \gamma^2} G_{3,0}^{1,3} \left( \frac{k^2 d^2}{\beta^2 \gamma^2} \sec^2 \psi_{0,1,1}^{3/2} \right),
\]  
(S17c)

\[
\mathcal{G}_p(\beta, k_d) = \frac{\sqrt{\pi}}{4} \int_{-\pi/2}^{\pi/2} \frac{d\psi}{\beta^2} G_{3,0}^{1,3} \left( \frac{k^2 d^2}{\beta^2 \gamma^2} \sec^2 \psi_{0,0,2}^{3/2} \right).
\]  
(S17d)

Here, \( k_x = \omega/c\beta, \kappa_x = \omega/c\gamma, \) and \( \alpha = e^2/4\pi \epsilon_0 \hbar c. \)
4. MAXIMUM OF SMITH–PURCELL RADIATION LIMIT

In Fig. 2, we discuss the limit of Smith–Purcell radiation at a given wavelength as a function of electron velocity. The electron velocity at which the limit of Smith–Purcell radiation achieves maximum corresponds to the zero of its derivative to velocity. In Eq. (4), since the integrand \( \kappa_4^4 K_0^2(\kappa \rho) + \kappa_5^2 k^2 K_1^2(\kappa \rho) \) is continuous and differentiable, based on the Lagrange’s mean value theorem, there must exists some \( \rho_0 \) such that \( \int_V \kappa_4^4 K_0^2(\kappa \rho) + \kappa_5^2 k^2 K_1^2(\kappa \rho) \, dV = \left[ \kappa_4^4 K_0^2(\kappa \rho_0) + \kappa_5^2 k^2 K_1^2(\kappa \rho_0) \right] V \). Therefore,

\[
\frac{d\Gamma(\omega)}{d\beta} \propto \frac{d\Gamma(\omega)}{d\kappa_4} \frac{d\Gamma(\omega)}{d\kappa_5} \frac{d\Gamma(\omega)}{d\rho}
\]

\[
\propto 4\kappa_4^2 K_0^2(\kappa \rho_0) - (3\kappa_4^4 \rho_0 + \kappa_5^2 k^2 \rho_0) K_0(\kappa \rho_0) K_1(\kappa \rho_0)
\]

\[
+ (4\kappa_4^3 + 2\kappa_5^2 k^2) K_1^2(\kappa \rho_0) - \rho_0 \kappa_4^2 (\kappa_5^2 + k^2) K_1(\kappa \rho_0) K_2(\kappa \rho_0).
\]

Figure S2. Plot of \( \frac{dN(\omega)}{d\omega \, d\beta} \bigg|_{\rho_0=2d} \). There is a nodal line of zero derivative that coincides with the limit maximum in Fig. 2.

Fig. S2 is calculated using Eq. S18 where a nodal line of zero derivative appears and coincide with the limit maximum shown in Fig. 2(a), which is consistent with our prediction of optimal velocities as a function of \( kd \).

5. LIMIT ASYMPTOTICS

For the asymptotic behavior of the limit, here we consider four scenarios: electrons in the near field \( (kd \rightarrow 0) \), electrons in the far field \( (kd \rightarrow \infty) \), extreme nonrealistic electrons \( (v \rightarrow 0) \), and
extreme relativistic electrons \((v \rightarrow c)\). In this section we only consider the three-dimensional problem \[\text{Eq. (4)}\].

First, we consider near field \(kd \rightarrow 0\). We also assume the electron speed is intermediate so neither \(\beta \rightarrow 0\) (extremely slow) nor \(\gamma \rightarrow \infty\) (extremely fast), which we will discuss later. In the expression of the general limit \[\text{Eq. (4)}\], there are two terms in the integrand where the first term (containing \(K_0\)) is the contribution from the longitudinal polarization \(E_x\) and the second term (containing \(K_1\)) is the contribution from the transverse polarization \(E_\rho\). The hyperbolic Bessel functions \(K_\nu\) in these two terms has the same argument \(\kappa_\rho \rho = k \rho / \beta \gamma\), which also approaches zero for \(\rho \geq d\). Both \(K_0(\kappa_\rho \rho)\) and \(K_1(\kappa_\rho \rho)\) diverge when \(\kappa_\rho \rho \rightarrow 0\) but at different divergence rates \[\text{Eq. (4)}\]:

\[
\lim_{\rho \to 0} K_0(\kappa_\rho \rho) \sim -\ln(\kappa_\rho \rho / 2) - \gamma_0, \quad (S19a)
\]

\[
\lim_{\rho \to 0} K_1(\kappa_\rho \rho) \sim \frac{1}{\kappa_\rho \rho}, \quad (S19b)
\]

where \(\gamma_0\) is the Euler–Mascheroni constant. Therefore, \(K_1(\kappa_\rho \rho) \gg K_0(\kappa_\rho \rho)\) when \(\kappa_\rho \rho \rightarrow 0\) and \(E_\rho\) has the major contribution to the limit.

Second, we consider electron beams in the far field \(kd \rightarrow \infty\):

\[
\lim_{\rho \to \infty} K_0(\kappa_\rho \rho) \sim \sqrt{\frac{\pi}{2 \kappa_\rho \rho}} e^{-\kappa_\rho \rho} \left[1 - \frac{1}{8 \kappa_\rho \rho} + O(\kappa_\rho^2 \rho^2)\right], \quad (S20a)
\]

\[
\lim_{\rho \to \infty} K_1(\kappa_\rho \rho) \sim \sqrt{\frac{\pi}{2 \kappa_\rho \rho}} e^{-\kappa_\rho \rho} \left[1 + \frac{3}{8 \kappa_\rho \rho} + O(\kappa_\rho^2 \rho^2)\right]. \quad (S20b)
\]

Therefore, both \(E_x\)–limit and \(E_\rho\)–limit decay exponentially at the same rate and \(E_\rho\)–limit remains be higher.

Third, we consider asymptotic behavior of the limit when the electrons are extremely nonrelativistic \((\beta \rightarrow 0)\). In this limit, we have \(\lim_{\beta \to 0} \kappa_\rho = k_v \rightarrow \infty\). Thus in \[\text{Eq. (4)}\]

\[
\lim_{\kappa_\rho \to \infty} \kappa_\rho^2 K_0(\kappa_\rho \rho) \sim \sqrt{\frac{\pi}{2 \kappa_\rho \rho}} \kappa_\rho^2 e^{-\kappa_\rho \rho} \left[1 - \frac{1}{8 \kappa_\rho \rho} + O(\kappa_\rho^2 \rho^2)\right] = 0, \quad (S21a)
\]

\[
\lim_{\kappa_\rho \to \infty} \kappa_\rho \kappa_v K_1(\kappa_\rho \rho) \sim \sqrt{\frac{\pi}{2 \kappa_\rho \rho}} \kappa_\rho^2 e^{-\kappa_\rho \rho} \left[1 + \frac{3}{8 \kappa_\rho \rho} + O(\kappa_\rho^2 \rho^2)\right] = 0, \quad (S21b)
\]

which is consistent with the fact that static charges do not generate radiation. Our computational verification is shown in Fig. 2(b) and (c) where both the limit and numerical results approach zero as \(\beta \rightarrow 0\) for either small or large separations (whether slow or fast electrons are preferred) between the electron beams and the structure.
Last, we consider the limit behavior when the electrons are extremely relativistic, where 
\[ \lim_{\beta \to 1} \kappa_{\rho} = \sqrt{\omega^2 / v^2 - \omega^2 / c^2} = 0: \]

\[ \lim_{\kappa_{\rho} \to 0} \kappa_{\rho}^2 K_0(\kappa_{\rho} \rho) \sim \kappa_{\rho}^2 \left[ -\ln(\kappa_{\rho} \rho / 2) - \gamma_0 \right] = 0, \quad (S22a) \]

\[ \lim_{\kappa_{\rho} \to 0} \kappa_{\rho} k_v K_1(\kappa_{\rho} \rho) \sim \kappa_{\rho} k_v / \kappa_{\rho} \rho = k_v / \rho. \quad (S22b) \]

Therefore, in this limit, \( E_x \) contribution vanishes but \( E_\rho \) remains finite. The entire problem becomes equivalent to a plane-wave scattering problem since the incident field is purely transverse.

6. PENETRATING ELECTRON TRAJECTORIES

In the main text, we discuss electron trajectories near photonic structures. For penetrating electron trajectories—that is, when the electron trajectory \( \mathbf{r}_e(t) \) intersects \( \chi(\mathbf{r}) \neq 0 \) regions—a subtlety arises: the limit, Eq. (3), then apparently diverges even in lossy materials \( \text{Im} \chi \neq 0 \). In specific terms, the norm-squared incident field \( E_{\text{inc}} \) is non-integrable over the electron trajectory, that is

\[ \int_V dV |E_{\text{inc}}(\mathbf{r})|^2 \sim \int_V dV |\hat{x} \kappa_{\rho} \ln(\kappa_{\rho} \rho + \hat{\rho} \gamma \rho^{-1})|^2 \]

diverges if \( V \) includes regions where \( \rho = 0 \). Here, we discuss the regularization of this divergence with emphasis on the implications to electron energy loss spectroscopy (EELS).

Though at first sight disconcerting, the divergence is not a surprise: the direct calculation of the EEL spectrum, \( \Gamma(\rho, \omega) = \frac{e^2}{\pi \hbar v^2} \text{Re} \int_{-\infty}^{\infty} dx E_x(\rho + x \hat{x}, \omega) e^{-ik_v z} \), is also divergent for penetrating trajectories when \( \text{Im} \chi \neq 0 \). For an extended bulk material, of permittivity \( \epsilon = 1 + \chi \), the EEL spectrum (per unit length \( L \)) can be evaluated from the momentum-space representation of the total field (to be introduced shortly), yielding [2]:

\[ \Gamma_{\text{EELS}}(\omega) = \frac{e^2 L}{\pi \hbar v^2} \text{Im} \left[ \left( \frac{\nu}{c} - \frac{1}{\epsilon} \right) \ln \left( \frac{q_c^2 + k_v^2 - \epsilon k_v^2}{k_v^2 - \epsilon k_v^2} \right) \right]. \quad (S23) \]

The denominator of the logarithm describes the emergence of Cherenkov losses for \( \nu > c/\epsilon \) and is finite—in contrast, the numerator, which describes EEL due to material loss, diverges logarithmically in a momentum cut-off \( q_c \). Of course, the divergence is merely an artifact of an idealized description of the system—several physical and practical considerations impose natural momentum cut-offs, e.g.:

**Collection angle:** The collection semi-angle of the microscope’s spectrometer \( \varphi \) restricts momentum transfer collection to in-plane momenta \( q_\rho < q_c \), with \( \hbar q_c = m_e v \sin \varphi \approx m_e v \varphi \). At typ-
ical collection semi-angles and acceleration voltages—say, \( \varphi = 10 \) mrad and 100 keV—which sets a cut-off at \( \hbar q_c \approx 2.8 \times 10^3 \text{ eV}/c \), or equivalently, a spatial spread \( 1/q_c \sim 1 \text{ Å} \).

**Nonlocality:** Nonlocality effectively suppresses the dielectric response to large-momentum plane-wave components, i.e., \( \epsilon(q, \omega) \rightarrow 1 \) for \( q \gg 1/a \) (lattice constant \( a \)). The free-electron response is quenched even earlier, at a threshold set by the Thomas–Fermi momentum.

**Electron spread:** The spread, \( \Delta R \), of the electron’s in-plane density imposes a cut-off \( q_c \sim 1/\Delta R \).

To summarize; the divergence of the limit for penetrating trajectories is simply the mirror of the divergence of the direct calculation. Accordingly, the divergence’s remedy is also mirrored: the limit is regularized upon introducing a momentum cut-off in the electron’s (incident) field \( E_{\text{inc}} \).

Denoting this regularized field \( E_{\text{inc}, q_c} \), we next verify that this field is indeed regular as \( \rho \rightarrow 0 \). Coincidentally, this also outlines the derivation of the conventional, non-regularized field [Eq. (2)].

The derivation proceeds as follows: in momentum-frequency space, the electron charge density \( \rho(\mathbf{r}, t) = -e\delta(\mathbf{r} - \mathbf{v}t) \) equals \( \rho(\mathbf{q}, \omega) = -2\pi e\delta(\omega - \mathbf{q} \cdot \mathbf{v}) \) and is accompanied by a current density \( \mathbf{J}(\mathbf{q}, \omega) = -2\pi e\mathbf{v}\delta(\omega - \mathbf{q} \cdot \mathbf{v}) \). Jointly with Maxwell’s equations, in the form of the wave-equation \( (q^2 - \epsilon k^2)E_{\text{inc}} = i\epsilon_0^{-1}(k/c - \mathbf{q}/\epsilon) \), this gives the associated electric field’s \( (\mathbf{q}, \omega) \)-representation:

\[
E_{\text{inc}}(\mathbf{q}, \omega) = -\frac{2\pi i e k v/c - \mathbf{q}/\epsilon}{q^2 - \epsilon k^2} \delta(\omega - \mathbf{q} \cdot \mathbf{v}).
\]

(S24)

An inverse transform then yields the \( (\mathbf{r}, \omega) \)-representation (specializing to \( \mathbf{v} = v\hat{x} \) and \( \epsilon = 1 \)):

\[
E_{\text{inc}, q_c}(\mathbf{r}, \omega) = -\frac{2\pi i e}{\epsilon_0} \int_{|\mathbf{q}| < q_c} \frac{d^3 \mathbf{q}}{(2\pi)^3} \frac{k v/c - \mathbf{q}}{q^2 - k^2} \delta(\omega - \mathbf{q} \cdot \mathbf{v}) e^{i\mathbf{q} \cdot \mathbf{r}}
\]

\[
= -\frac{ie}{\epsilon_0 v} e^{ik_x x} \int_{|\mathbf{q}| < q_c} \frac{d^2 \mathbf{q}_\perp}{(2\pi)^2} \frac{2q v}{q^2 + k_x^2 - k^2} (k v/c - k_x) \hat{x} - \mathbf{q}_\perp e^{i\mathbf{q}_\perp \cdot \mathbf{r}}
\]

\[
= \frac{ie}{2\pi \epsilon_0 v} e^{ik_x x} \left[ \frac{k_v}{\gamma^2} \int_0^{q_c} dq \frac{q \rho J_0(q \rho)}{q^2 + k_x^2 - k^2} \hat{x} + i \int_0^{q_c} dq \frac{q^2 J_1(q \rho)}{q^2 + k_x^2 - k^2} \hat{\rho} \right],
\]

(S25)

reproducing Eq. (2) as \( q_c \rightarrow \infty \) (we remind that \( \kappa_0 \approx k_v / \gamma \)). Written in terms of the transverse and longitudinal parts introduced in the above, \( L_{q_c} \) and \( T_{q_c} \), the regularized version of Eq. (3) reads

\[
P_\tau(\omega) \leq \frac{e^2 \omega^2 \xi^2}{16 \pi^3 \epsilon_0 v^2} \int dV \frac{|\chi|^2}{\text{Im} \chi} \left(L_{q_c}^2 + T_{q_c}^2\right).
\]

(S26)

To demonstrate the limits’ finiteness, we require the small-\( \rho \) behavior of \( L_{q_c} \) and \( T_{q_c} \) at finite \( q_c \).
Since $q_c$ is large, much larger than $\kappa_\rho$, this is straightforward—particularly for $T_{q_c}$:

$$T_{q_c} \doteq \int_{0}^{q_c} \frac{q^2_{\rho} J_1(q_{\rho}\rho)}{q^2_{\rho} + \kappa^2_{\rho}} = \int_{0}^{\infty} \frac{q^2_{\rho} J_1(q_{\rho}\rho)}{q^2_{\rho} + \kappa^2_{\rho}} - \int_{q_c}^{\infty} \frac{q^2_{\rho} J_1(q_{\rho}\rho)}{q^2_{\rho} + \kappa^2_{\rho}} = \kappa_{\rho} K_1(\kappa_{\rho}\rho) - \int_{q_c}^{\infty} \frac{q^2_{\rho} J_1(q_{\rho}\rho)}{q^2_{\rho} + \kappa^2_{\rho}} \frac{J_0(q_{\rho}\rho)}{\rho} = \kappa_{\rho} K_1(\kappa_{\rho}\rho) - \frac{J_0(q_{\rho}\rho)}{\rho}.$$  (S27)

The small-$\rho$ behavior then follows from the small-argument asymptotics of the Bessel functions

$$T_{q_c} \approx \frac{\kappa_{\rho}}{\gamma} \int_{0}^{q_c} \frac{q^2_{\rho} J_0(q_{\rho}\rho)}{q^2_{\rho} + \kappa^2_{\rho}} \sim \frac{\kappa_{\rho}}{\gamma} \left[ \int_{0}^{\infty} \frac{q^2_{\rho} J_0(q_{\rho}\rho)}{q^2_{\rho} + \kappa^2_{\rho}} - \int_{q_c}^{\infty} \frac{q^2_{\rho} J_0(q_{\rho}\rho)}{q^2_{\rho} + \kappa^2_{\rho}} \right] = \frac{\kappa_{\rho}}{\gamma} \left\{ K_0(\kappa_{\rho}\rho) + \ln \frac{1}{\gamma} \left( \frac{1}{2} q_c \rho + \gamma_{\text{em}} - \frac{1}{2} \left( \frac{1}{2} q_c \rho \right)^2 \right) \right\},$$

(S28)

where $2F_3$ is a generalized hypergeometric function with the asymptotic behavior $1 - O[(q_c\rho)^2]$. The small-$\rho$ behavior again follows from the Bessel function asymptotics $K_0(x) = -\ln \frac{1}{\gamma} x - \gamma_{\text{em}} + O(x^2 \ln x)$, such that:

$$L_{q_c} \approx \frac{\kappa_{\rho}}{\gamma} \ln \frac{q_c}{\kappa_{\rho}}, \quad \text{for } \rho, q_c \ll \kappa_{\rho}^{-1}.$$  (S29)

Thus, the longitudinal contribution $L_{q_c}$ tends to a finite, nonzero value $\propto \ln q_c/\kappa_{\rho}$ as $\rho \to 0$; this is also the maximum of $L_{q_c}$.

Equations (S28) and (S29) demonstrate that the $\rho = 0$ singularity of the incident field is regularized for finite cut-off momenta $q_c$. This ensures that both direct calculations and limits similarly yield finite, regularized values, with bulk contributions dependent on the cut-off momentum.

7. EXPERIMENTAL METHODS AND DATA ANALYSIS

We are able to obtain the absolute intensity of Smith–Purcell radiation by implementing a calibration measurement using a broadband (visible and near infrared) calibrated source (Avalight-HAL-CAL). The experimental setup for calibration is shown in Fig. S3. All the optics remain the
same as Fig. 3(a) except that we replace the SEM with the calibration source. The spectral intensity $S_0(\omega)$ of the calibrated source is already known from the manufacturer. Passing through all the optics, the radiation from the calibrated source enters the spectrometer and generates a signal count $C_0(\omega)$. 

With $S_0(\omega)$ and $C_0(\omega)$, we are able to gauge Smith–Purcell radiation intensity $S(\omega)$ by reading the corresponding signal count $C(\omega)$. The relation is given by

$$\frac{S_0(\omega)}{C_0(\omega)} = \frac{S(\omega)}{C(\omega)}.$$  

(S31)

This relation is valid for two reasons. First, the generated photons into the sample substrate is negligibly small compared to the total radiation (see Fig. S4). Second, the optics and spectrometer configurations remain unchanged for Smith–Purcell radiation measurement and calibration measurement. This approach allows us to obtain the absolute radiation intensity of the collected Smith–Purcell radiation, without knowing the loss functions of each individual optical elements or the quantum efficiencies and EM gains of the spectrometer at each wavelength, since all these factors will cancel out if inserted into Eq. (S31).

To calculate the number of photons generated per electron, measurement of the current from the SEM is necessary. The currents are measured using a picoammeter connected to a built-in Faraday cup inside the SEM chamber. The measured currents are shown in Fig. S5(a).

For comparisons with the analytical limits, we also need to evaluate the number of unit cells $N_{uc}$ of interaction and consider the beam diameters (spatial spread) of the electron beams. We estimate the electron beam diameter $D$ with the equation [7]

$$D^2 = D_0^2 + D_d^2 + D_s^2 + D_c^2 = \left[C_0^2 + (0.6\lambda)^2\right] \alpha_p^{-2} + \frac{C_s^2 \alpha_p^6}{4} + \left[C_c \frac{\Delta E}{E}\right]^2 \alpha_p^2.$$  

(S32)

Here $D_0$ is the aberration-free Gaussian probe diameter, $D_d$ corresponds to aperture diffraction, $D_s$ corresponds to spherical aberration, and $D_c$ corresponds to chromatic aberration. Our SEM
Figure S4. Fraction of the generated photons into the substrate for different accelerating energies at normal emission angle ($\lambda = a/\beta$) for the first-order Smith–Purcell radiation.

uses a tungsten thermionic cathode, for the energy regime (10–20 keV) we use, $D_d$ and $D_c$ are negligible [8]

$$D^2 \approx D_0^2 + D_s^2 = C_0^2 \alpha_p^{-2} + \frac{C_s^2 \alpha_p^6}{4},$$

(S33)

where

$$C_0 = \sqrt{4I/b\pi^2},$$

(S34)

$b$ is the electron gun brightness, $I$ is the probe current, $\alpha_p$ is the convergence semi-angle of the electron beam, and $C_s$ is the spherical aberration coefficient. For the brightness $b$ of the source, we choose $1 \times 10^5$ A/cm$^2$/sr for the acceleration energy 20 keV (typical value for a tungsten source [7–9]) and scale it linearly [7–9] for other voltages. The focal length (working distance) of our SEM is 28 mm, which corresponds to a spherical aberration coefficient $C_s \approx 300$ mm [8, 9].

Figure S5. (a) Measured current of the experiment. (b) Schematic of the model to evaluate the interaction length of the electron beam with the structure. (c) Electron structure separations $d$ obtained from the model (dots) and their polynomial fitting (curve; the 20 kV outlier data point dropped from fitting) for calculating theoretical upper limits.
For each measurement, we adjusted the SEM to achieve the smallest possible beam diameter. In theory, this corresponds to \( D_{\text{min}} = (4/3)^{3/8}(C_0^3 C_s)^{1/4} \) for the optimal convergence semi-angle \( \alpha_{\text{opt}} = (4/3)^{1/8}(C_0/C_s)^{1/4} \) [derived from Eq. (S33)].

In our experiment, the electron beams grazingly impinges onto the sample at a nonzero angle of \( \theta = 1.5^\circ \), which leads to a finite number of unit cells where electrons strongly interact with the structure such that the radiation contribution from other areas are negligible. The backscattering coefficient \( \eta \) of the SEM can be generally estimated as [7]

\[
\eta = 1/(1 + \sin \theta)^p,
\]

where \( p = 9/\sqrt{Z} \) and \( Z \) is the atomic number. In our case, \( \theta = 1.5^\circ \) and \( Z = 79 \) (Au), and thus \( \eta \approx 0.974 \), meaning that most electrons get elastically scattered and maintain their initial momenta, which correspond to the scenario shown in Fig. S5(b). The highlighted rectangle is treated as the region where electrons strongly interact with the structure. The number of unit cells is consequently determined via the length of the interaction region \( N_{\text{uc}} = L/a = 2D/a \sin \theta \). After obtaining \( N_{\text{uc}} \), the measured radiation spectral density \( S(\omega) \) can be translated into emission probability per electron per frequency per unit propagation length

\[
\frac{d\Gamma_{\text{expt}}(\omega)}{dx} = \frac{eS(\omega)}{\hbar \omega IN_{\text{uc}}a},
\]

which produces the measured emission probability shown in Fig. 3(d).

On the theory side, the upper limit in Fig. 3(d) is calculated for Smith–Purcell radiation at the surface-normal emission angle (i.e., \( \beta = a/\lambda \)). The limit is evaluated at the center of the interaction region with separation \( d = H/2 = D \tan \theta/4 \sin \theta \) [see Fig. S5(c)] by generalizing Eq. (6). The generalization of Eq. (6), analogous to the expression of Eq. (4), is straightforward for the inhomogeneous Au-Si grating sample: move \( |\chi|^2/\text{Im}\chi \) into the integrand, and account for different materials:

\[
\Gamma_{\tau}(\omega) \leq \frac{\alpha \xi c}{2\pi \omega^2} \sum_{\text{mat}} \int_{V_{\text{mat}}} \frac{|\chi_{\text{mat}}|^2}{\text{Im}\chi_{\text{mat}}} \left[ k_p^4 K_0^2(k_p\rho) + k_p^2 k_y^2 K_1^2(k_p\rho) \right] dV,
\]

where \( V_{\text{mat}} \) and \( \chi_{\text{mat}} \) are the occupied volume and susceptibilities of the materials (mat \( \in \{\text{Si, Au}\} \)).

8. NEAR–INFRARED SMITH–PURCELL RADIATION EXPERIMENT

We also conduct near–infrared experiment to further confirm our theory with the same experimental setup and a near–infrared spectrometer. A one-dimensional grating (Au-covered patterned-
Si, see Fig. S6 inset; LightSmyth Technologies) with a longer periodicity (≈ 272 nm) is used such that the Smith–Purcell radiation moves to near–infrared.

Adopting the same methods of data acquisition, calibration, and analysis [as those of our initial experiment in the visible (as described in Supplementary 7)], we are able to obtain the absolute emission probabilities for the near–infrared Smith–Purcell radiation. The new experimental results are shown in Fig. S6, where the envelope lineshape of the emission spectra again follows our theoretical prediction. The measured currents and the calculated electron structure separations are shown in Fig. S7.

Figure S6. Smith–Purcell radiation observed in the near–infrared regime and the comparison with the upper limit theory.

Figure S7. (a) Measured current of the near–infrared experiment. (b) Electron structure separations $d$ obtained from the model (dots; see Supplementary 7) and their polynomial fitting (curve) for calculating theoretical upper limits.

In addition to the agreement between our theory and each of the experiment, the comparison between the visible and the infrared experiment gives rise to interesting observations that further
confirm our theory. Two key observations can be made from the comparison. First, the absolute emission probabilities increase by about two orders of magnitude from the visible to the near-infrared regime—consistent with the same order of increase in the material factor of Au [see Fig. 1(b)], which confirm the material factor dependence explicitly. Second, although the two experiments are both in the fast-electron-efficient regime, the measured emission probabilities feature a peak for the visible experiment, while exhibit monotonic decrease for smaller electron energies (except for a small increase between 17 keV to 16 keV) for the near–infrared experiment. Such a difference arises because the material response is much less dispersive in the near–infrared, which implicitly corroborates the functional impact-parameter dependence within our upper limit.

9. COMPLEMENTARY EVIDENCE FOR THE SLOW-ELECTRON-EFFICIENT PREDICTION

In the main text, we predict that slow electrons radiate more strongly than relativistic ones at subwavelength separation ($kd \ll 1$) with structures. We also provide numerical evidence for this prediction [Fig. 2(b-c)]. In this section, we discuss a complementary supporting evidence for our slow-electron-efficient prediction based on data extracted from a recent work [10] that reports an integrated Cherenkov radiator using hyperbolic metamaterials (Au/SiO$_2$ layered stack). The electron-structure separation is reduced by integrating the electron field emitter on the chip. Ref. [10] reports the output power $P_{\text{out}}$ of the device as a function of anode-cathode currents $I_{\text{ac}}$ and electron energies for fixed radiation wavelengths (centered at ≈780 nm; see Fig. 2 in Ref. [10]). These data allow us to extract the experimental emission probabilities $\frac{d\Gamma(\omega)}{dx} \propto P_{\text{out}}/I_{\text{ac}}$, since the reported shapes of the radiation spectra are similar for various electron energies (i.e., almost constant emission bandwidth for various electron energies).

The probabilities are therefore shown in Fig. S8, where data reported with electron energies are shown with extra red circles. Data points without electron energies reported in [10] are interpreted as linearly-interpolated electron energies. As we explicitly show in Fig. 2, the lineshape of emission probabilities versus electron velocity contains one-to-one correspondence with $kd$. Hence, the experimental lineshape can be fitted with the shape-independent upper limit [Eq. (5a)] to extract the electron-structure separation $d$ (being the only free parameter to compare with the reported value):

$$\frac{\Gamma_{\text{rad}}(\omega)}{dx} \propto \frac{|\chi_{\text{eff}}|^2}{\Im \chi_{\text{eff}} \beta^2} \left[ (\kappa_p d)K_0(\kappa_p d)K_1(\kappa_p d) \right],$$

(S38)
Figure S8. Extra external supporting evidence for the slow-electron-efficient prediction with small electron energies (0.25–1.4 keV), which complements our experiments using electrons with higher energies (10–20 keV). Extracted emission probabilities from [10] and the optimal lineshape fitting (solid lines with $kd \approx 0.16$) using the upper limit. The auxiliary suboptimal fittings (dashed and dotted curves) demonstrate the uniqueness of the extracted $kd$ value. All red curves fit data points with electron energies reported, while all black curves fit all data points assuming linearly interpolated electron energies (see Fig. 2 in Ref. [10]).

where $\chi_{\text{eff}}(\chi_{\text{Au}}, \chi_{\text{SiO}_2}, \beta)$ is the effective susceptibility of the metamaterial, a function of the composite material permittivities and the longitudinal wavevector (i.e., the electron velocity), is explicitly determined from the standard effective medium theory (see Supplementary Information of Ref. [10] S1–S3 sections). We obtain two primary fitting results in Fig. S8, where $\hat{d}_1 = 20.0 \pm 2.3$ nm for the solid black curve (fits all data) and $\hat{d}_2 = 19.8 \pm 6.7$ nm for the solid red curve (only fits the data with electron energies reported). The estimated $kd \approx 0.16$ corresponds to an optimal nonrelativistic electron velocity $\beta_{\text{opt}} \approx 0.08$ (see Fig. S8). We emphasize that although the only available experimental data are below the predicted $\beta_{\text{opt}}$, they are sufficient for us to unambiguously extract the $kd$ value for the experiment. The uniqueness of the $kd$ value is supported by the two auxiliary suboptimal fittings (dashed and dotted curves) with either reduced or increased $kd$ values.

For comparison, the reported separation from the field emitter to the structure is 40 nm. Since the electron beam in the reported device is still not theoretically ideal (not a delta function in space; instead, with nonzero beam diameters), we consider the estimates $\hat{d}_1$ and $\hat{d}_2$, from our upper limit theory, as good agreement with the realistic structural parameter. Hence, such an agreement serves
as complimentary support for our prediction about the slow-electron-efficient regime.

10. UPPER LIMIT IN TWO DIMENSIONS

The limits can be derived in both the three-dimensional or the two-dimensional case. For completeness, here we also derive the limit in the two-dimensional case, which correspond to sheet electron beams that are assumed in Fig. 4(f).

We consider an electron sheet beam in the \((x, z)\) plane with charge density being one electron per nanometer, i.e., \(q = 1.6 \times 10^{-19} \text{ C/nm} \) [consistent with our unit for probability in two dimensions \(\frac{\rho_r}{\delta x \delta y} / \hbar \text{(eV}^{-1}\text{nm}^{-2})\)]. Precisely, the probability is invariant of the choice of the transverse \((y)\) length scale, as long as the length scale is in the same unit for both the source current density and the probability. Here the length scale is chosen as nanometer for both of the quantities.

The source current density in the time domain can be written as \(J(r, t) = qv\delta(z - z_0)\delta(x - vt)\hat{x}\).

In the frequency domain, the current density is given by

\[
J(r, \omega) = q\delta(z - z_0)e^{ik_v x}\hat{x},
\]

The induced fields are

\[
\begin{align*}
H(r, \omega) &= -\frac{q}{2}e^{ik_v x - \kappa_p(z - z_0)}\hat{y} \\
E(r, \omega) &= \frac{q}{2\omega\epsilon_0}(k_v\hat{z} - i\kappa_p\hat{x})e^{ik_v x - \kappa_p(z - z_0)}
\end{align*}
\]

for \(z > z_0\) and

\[
\begin{align*}
H(r, \omega) &= \frac{q}{2}e^{ik_v x + \kappa_p(z - z_0)}\hat{y} \\
E(r, \omega) &= -\frac{q}{2\omega\epsilon_0}(k_v\hat{z} + i\kappa_p\hat{x})e^{ik_v x + \kappa_p(z - z_0)}
\end{align*}
\]

for \(z < z_0\), where \(\epsilon_0\) is the vacuum permittivity, and \(\kappa_p\) also defined as \(\kappa_p = \sqrt{k_v^2 - k^2}\), same as the main text. where \(k = \omega/c\) is the light wavevector.

Insert Eq. \((S41b)\) into Eq. \((4)\), we obtain the limit in two dimensions

\[
\frac{d\Gamma_r(\omega)}{dy} \leq \frac{|\chi|^2}{\text{Im} \chi} \frac{q^2 \xi_r (k_v^2 + \kappa_p^2)}{32\hbar\epsilon_0 \omega^2} \int_S e^{-2\kappa_p|z - z_0|} dS,
\]

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where $S$ is the area defined by the profile of the structure.

As in the main text, we also consider a concrete example: Smith–Purcell radiation from a rectangular grating with filling factor $\Lambda$. Applied the rectangular profile to Eq. (S42), the radiated photon per frequency per electron per unit area is bounded by

$$\frac{\text{d}^2 \Gamma_r(\omega)}{\text{d}x \text{d}y} \leq \frac{|\chi|^2}{\text{Im} \chi} \frac{\Lambda q^2 \xi_r(k_x^2 + k_y^2)}{64 \hbar \epsilon_0 \kappa_x \omega^2} e^{-2 \kappa_x d},$$

where $d$ is the distance between the electron and the grating.

11. FREE ELECTRON RADIATION NEAR A BOUND STATE IN THE CONTINUUM

In photonic systems, modes below the light cone are guided modes, while modes above the light cone are typically resonances with finite lifetime. In contrast to guided modes or resonances, a bound state in the continuum is a perfectly confined modes with infinite lifetime embedded in the radiation continuum (above light cone) [11].

These properties can be used to distinguish whether an optical resonance is a BIC or not. The mode profile in Fig. 4(c) corresponds to a BIC because 1) it is obviously within the continuum [see Fig. 4(a)]; 2) it possesses infinite lifetime without external radiation (no outgoing oscillatory radiation in its eigenmode profile), as shown in Fig. S9(a).

![Figure S9. Difference of a BIC and a resonance on the TE$_1$ band of Fig. 4(a).](image)

(a) The mode profile of a BIC decays exponentially, giving rise to infinite lifetime; (b) The mode profile of a resonance contains oscillatory radiation into the far field, leading to finite lifetime.
Next we explain the Smith-Purcell radiation enhancement near a BIC. We write down the temporal coupled mode theory \[\text{(12)}\] for the coupling process

\[
\frac{da}{dt} = -i\omega_0 a - a \left( \frac{1}{\tau_{bg}} + \frac{1}{\tau_r} \right) + \sqrt{2 \tau_r} s_+,
\]

where \(a\) is the mode amplitude inside the resonances, \(1/\tau_{bg}\) is the background coupling rate (such as material absorption and scattering loss due to fabrication imperfections), \(1/\tau_r\) is the resonant coupling rate (for BIC, \(\tau_r = \infty\)), \(s_+\) is the wave amplitude carried by the electron towards the resonances. Solving Eq. \((S44)\) for \(a\) we have

\[
\left| \frac{a}{s_+} \right|^2 = \frac{2/\tau_e}{(\omega - \omega_0)^2 + \left( \frac{1}{\tau_{bg}} + \frac{1}{\tau_r} \right)^2},
\]

Assuming on-resonance condition \((\omega = \omega_0, \text{electron line and photonic bands intersects})\), to maximize the resonance amplitude for a given fixed \(s_+\), we have

\[
\left| \frac{a}{s_+} \right|^2 = \frac{2/\tau_r}{\left( \frac{1}{\tau_{bg}} + \frac{1}{\tau_r} \right)^2} \propto \frac{Q_{\text{tot}}^2}{Q_r},
\]

where \(Q = \omega \tau/2\) for all channels and \(1/Q_{\text{tot}} = 1/Q_r + 1/Q_{bg}\). It is thus evident from Eq. \((S46)\) that the maximal resonance enhancement is achieved when \(Q_{bg} = Q_r\), (i.e., \(\tau_{bg} = \tau_r\)) which is the “Q-matching condition” we refer to in the manuscript.
As a result, the achievable radiation enhancement depends on the background radiation rate (see Fig. S10). In our example shown in Fig. 4(f) with material absorption taken into account, the maximal enhancement occurs at small offsets from the BIC with a $Q_r \approx 10^3 \sim 10^5$.

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