Transition radiation in metal-metal multilayer nanostructures as a medical source of hard x-ray radiation

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We show that a periodic metal-metal multilayer nanostructure can serve as an efficient source of hard x-ray transition radiation. Our research effort is aimed at developing an x-ray source for medical applications, which is based on using low-energy relativistic electrons. The approach toward choosing radiator-spacer couples for the generation of hard x-ray resonant transition radiation by few-MeV electrons traversing solid multilayer structures for the energies of interest to medicine (30–50 keV) changes dramatically compared with that for soft x-ray radiation. We show that one of the main factors in achieving the required resonant line is the absence of the contrast of the refractive indices between the spacer and the radiator at the far wings of the radiation line; for that purpose, the optimal spacer, as a rule, should have a higher atomic number than the radiator. Having experimental goals in mind, we have considered also the unwanted effects due to bremsstrahlung radiation, absorption and scattering of radiated photons, detector-related issues, and inhibited coherence of transition radiation due to random deviation of spacing between the layers. Choosing as a model example a Mo–Ag radiator-spacer pair of materials, we demonstrate that the x-ray transition radiation line can be well resolved with the use of spatial and frequency filtering.


I. INTRODUCTION

One of the promising medical applications of hard x rays is a so called dichromatic x-ray contrast diagnostics, whereby the absorption edge of a dye of interest (e.g., iodine) is “bracketed” by two (usually very broad) bands of radiation. The x-ray source for that application explored so far was synchrotron radiation of a large accelerator, which makes it impractical for wider use. To make it compact and practical, one needs to develop an x-ray source with narrow radiation lines of sufficient photon flux and much lower electron accelerator energy. An ideal way to go would be to employ the narrow-line resonant transition radiation (TR) of electrons traversing a multilayer solid-state nanostructure, proposed in Ref. 5. Since the main domain of interest are hard x rays beyond, e.g., 20 keV, the only effect that may provide any noticeable contrast of refractive index between the nanostructure layers is the narrow-line change of the refractive index in the close vicinity of absorption lines (so called atomic edges) of K, L, or M shells of one of the alternating layer elements (the so called radiator alternated by nonresonant spacers).

Our research effort is aimed at developing an x-ray source that would rely on the use of low-energy electrons, and it is based on the earlier work of two of us, which has theoretically demonstrated feasibility of generating intense x-ray transition radiation by few-MeV electrons traversing solid multilayer nanostructures. Generation of soft x rays via resonant TR in a multilayer submicron structure, albeit without the use of the dielectric contrast near the atomic photoabsorption edges, has been experimentally demonstrated in Ref. 6. The most dramatic difference of the results (see also Ref. 7) from the main body of works on x-ray TR (Ref. 8) is the possibility to use electron (E) beams of 5–10 MeV energy to generate x rays in 30–50 keV range, whereas the conventional technology, based on foil stacks, needs electron energies three to four orders of magnitude higher. As a result, widely available and relatively inexpensive industrial and medical electron accelerators could be used instead of electron synchrotrons of the national-facility kind, and thus, low-cost sources of x rays for numerous practical applications can be developed. Another method we employ, also based on results, is the utilization of resonances at inner-shell absorption edges of materials to narrow the bandwidth of the generated radiation and greatly enhance the intensity of the TR at each interface due to relatively large contrast of dielectric constants of the adjacent layers based on the huge jumps of absorption in the spectral vicinity to the inner-shell absorption edges.

The paper is organized as follows: In Sec. II we discuss the problem of choosing the radiator-spacer pairs. We formulate the requirements on the material parameters and compare with those for the materials suitable for generation of TR in the soft x-ray domain. Next, we provide expressions for calculation of the TR differential efficiency (Sec. III), based on which we choose specific pairs of materials and
study their radiation efficiency (Sec. IV). Comparison of the TR with the bremsstrahlung (BS) radiation, the main source of “noise” limiting efficiency of the metal-metal nanostructures as a source of hard x-ray TR, is provided in Sec. V. Section VI is devoted to discussing the measurement arrangement and detectors suitable for the proof-of-principle experiment we have planned. Then we discuss effects of the random layer width fluctuations (e.g., due to manufacturing inaccuracy) on the coherence of the TR; we also calculate precision tolerance of the nanostructure period allowing for substantial coherency gain in the output signal (Sec. VII).

II. SELECTION OF RADIATOR-SPACER PAIRS

The requirement of sufficiently hard x rays suggests relatively heavy atoms with respectively high $K$ shells as the choice for a radiator. Currently, the most used element for a dye is iodine; recent research results indicated that better results from the medical point of view can be achieved by using Gadolinium. Our preliminary search identified two “bracketing” radiator elements for iodine ($Z=53$, $E_K = 33 169$ keV), Te ($Z=52$, $E_K \approx 31 814$ keV) and Ba ($Z=56$, $E_K = 37 441$ keV), and two “bracketing” radiator elements for gadolinium ($Z=64$, $E_K = 50 239$ keV), Eu ($Z=63$, $E_K = 48 519$ keV) and Tb ($Z=65$, $E_K = 51 995$ keV).

Our original approach aimed mostly at soft x-ray domain, and it was based on choosing the layer of heavy atoms as a radiator with a chosen $K$-shell transition, and the layer of lighter atoms as a neutral spacer. However, in our detailed calculations we realized that approach while good for the soft x-ray domain does not produce desirable results in the hard x-ray domain. The major drawback is that in the 20–50 keV x-ray domain the TR spectrum with such pairs shows a spectral dip at the chosen $K$ shell, instead of a desired spectral peak (see top curve in Fig. 1). While this effect may be beneficial for certain applications, in our case it proved to be counterproductive.

The problem on how to break through this obstacle by finding an approach to the proper choice of the radiator-spacer pairs arose. Our search through the periodic table of elements resulted in locating a few candidates of spacers for any given radiator to produce strongly pronounced resonant peak of transition radiation with the contrast better than two orders of magnitude. Surprisingly enough, these spacer candidates as a rule are heavier than the radiator. One of the requirements on the candidates we imposed is that they have to be technologically viable and accessible, which narrows down the field of candidates, but still leaves at least two candidates for each radiator element. The very major factor in the TR excited by electrons passing though a layered structure is that it occurs because of the difference (contrast), $\Delta \varepsilon = \varepsilon_r - \varepsilon_s$, between the dielectric constants of the adjacent layers at the frequency $\omega$ (or photon energy $E_{ph} = h\omega$) of radiation, where the subscripts $s$ and $r$ denote spacer and radiator, respectively. The energy of the TR at the frequency $\omega$ is proportional to $|\Delta \varepsilon(\omega)|^2$. Here are the major requirements for the choice of materials.

- The radiator and spacer should have a largest possible $\Delta \varepsilon$ between them at the central frequency of the radiation line. In the domain of interest, the atomic absorption edges are due to $K$-shell ionization, and the radiator should comprise relatively heavy atoms, starting from Mo ($E_K = 20$ keV).
- At the same time, the radiator and spacer should have a very close dielectric constants at the wings of the radiation line, since otherwise, there will be a strong TR at the far wings, and the peak of radiation at the $K$-shell frequency will be very indistinct and of no use for the application in consideration.

Essentially, this is a signal-to-noise ratio problem; we should be seeking for the couples with $|\Delta \varepsilon_{\text{wings}}|^2 \ll |\Delta \varepsilon_{\text{center}}|^2$. This wing-matching condition immediately narrows down our options to both the spacer and radiator comprising heavy atoms, or more precisely, having high nuclei Coulomb charge $Z$. The latter condition is based on the following considerations. The lightest atom in the desirable domain is Mo, with nuclei Coulomb charge, and thus total number of electrons in a neutral atom, being $Z=42$.

The rest of the atoms in that domain have even higher $Z$. In the frequency domain of interest the dielectric constant of material could be written as $\varepsilon = 1 - \frac{2\delta}{\chi Z}$, with $0 < \delta < 1$. Away from the $K$ atomic absorption edge of the radiator yet reasonably close to it (within a few percents of the frequency), we have $\delta \approx f_1$ with $f_1 \sim Z$, where $0 < Z - f_1 \ll Z$ and $f_1$ is the real part of the so-called scattering factor. At the very edge of the atomic $K$ resonance, the largest possible deviation of $f_1$ from $Z$ would be about $\ln(\Gamma_K)$ where $\Gamma_K$ is the natural (i.e., due to radiation) dumping factor, and it is getting larger for heavier atoms. Thus, the largest difference $Z - f_1 \sim \ln(\Gamma_K)$ would be around 9 (in real terms it is usually lower than 8), with the relative contrast being $\sim 20\%$ or even lower for heavier atoms.

If one chooses now a spacer with much lighter atoms, its dielectric constant would be determined by $\delta_s \approx Z_s$, where $Z_s \ll Z$ is the total number of atomic electrons for the spacer atoms [for high frequencies near the atomic edge of a heavy radiator, we have $(f_1)_s = Z_s$]. Thus, the difference $\Delta \varepsilon = 2(\delta_r - \delta_s)$ will be almost fully determined by the heavy atoms of the radiator, with the spacer having very little impact, since $\delta_s \ll \delta_r$, and thus $\Delta \varepsilon \approx 2\delta_r$. Of course, $\delta$ depends also on

![Figure 1](https://example.com/image1.png)

FIG. 1. (Color online) The contrast of dielectric constants $|\Delta \varepsilon(\omega)|^2$ for Mo–Si, Mo–Ti, Mo–Ag, Mo–Pb, and Mo–Th (upper to lower curves, respectively).
the atomic density, but taking it into consideration does not change the matter significantly; in most of the cases it only further diminishes the contribution of the spacer atoms. Therefore, for all the practical purposes, the system with light spacer+heavy radiator would behave almost like vacuum+radiator, so that at the wings there will be strong background radiation with its intensity proportional to \(Z_i^2\), whereas at the atomic absorption edge of the radiator, it will be proportional to \((Z_i^2 - [\ln \Gamma_i])^2\), i.e., there will be a dip in the spectrum of radiation, rather than a resonant peak. As an example, we considered the technologically simplest pair, Mo–Si, which is well researched and made to order as extreme ultraviolet (EUV) mirrors: TR spectrum for such a pair would exhibit a dip near \(E_k\) of Mo (see upper curve in Fig. 1).

This explains why the “light spacer+heavy radiator” idea, good for the soft x-ray domain, fails to deliver an expected resonant line for the applications we are aiming at. Thus, for the hard x-ray domain one has to use the couples with the spacer being also a heavy element having \(Z_i\) fairly close to \(Z_s\), so that the “wing-match” condition would be satisfied and, therefore, the radiation at the wings would be quenched by having almost vanishing \(\Delta \varepsilon\). The rule of thumb for the acceptable formula becomes “heavy radiator +heavier spacer.”

### III. TRANSITION RADIATION EFFICIENCY OF A MULTILAYER STRUCTURE

Spatial intensity distribution of resonant TR is conical, with most of the intensity at each particular wavelength \(\lambda\) concentrating about a certain emission angle \(\theta\) measured from the electron velocity. Each TR spatial mode with a number \(r\) has a different resonant angle \(\theta_r\) related to the wavelength and to the electron velocity \(v\) by the resonant condition

\[
\frac{\varepsilon_1^{1/2}}{v} \cos \theta_r = \frac{c}{v} - \frac{r\lambda}{l}, \tag{1}
\]

where \(\varepsilon_1^{1/2} = (\sqrt{\varepsilon_1}l_1 + \sqrt{\varepsilon_2}l_2)/l\) is the mean refractive index of the medium, \(l_1\) and \(l_2\) are the thicknesses of different individual layers, \(l=l_1+l_2\) is the spatial period, and \(c\) is the velocity of light in vacuum. Equation (1) provides the condition for a constructive coherent interference of electromagnetic waves generated at different interfaces, at a distant point. Transition radiation differential efficiency in a multilayer structure, defined as the number of photons \(N\) per electron per unit solid angle \(\Omega\) per unit interval of the TR frequency, can be expressed as \[2,5,11-14\]

\[
\frac{d^2N}{d\Omega d\omega} = F_1 F_2 F_3, \tag{2}
\]

where \(F_1\) is the differential efficiency of a single interface, \(F_2\) reflects the contribution of a single plate (i.e., of the very first couple of adjacent interfaces), and \(F_3\) represents coherent summation of radiation from all the periods (i.e., couples of individual adjacent layers). For a strongly relativistic beam, i.e., when \(\beta = v/c \approx 1\) and the relativistic factor \(\gamma = 1 - \beta^2\) is high, \(\gamma^2 \gg 1\), the differential efficiency of a single interface is

\[
F_1 = \frac{\alpha (\Delta \varepsilon)^2}{\pi^2 \omega} |G|^2, \tag{3}
\]

where \(\Delta \varepsilon = \varepsilon_1 - \varepsilon_2\) is the difference between the dielectric constants of the structure layers, \(\alpha = 1/137\) is the fine structure constant, and \(G\) is a radiation pattern of a single interface given by

\[
G = -\frac{\theta}{(\gamma^2 + \theta^2)^{1/2}}, \quad \theta \ll 1, \tag{4}
\]

so that the angle of maximum radiation is \(\theta_{\text{max rad}}\) = \(1/(\gamma \sqrt{3})\).

The factor \(F_3\) in Eq. (2) is due to coherent summation of radiation generated at the very first couple of adjacent interfaces. If we choose an angle of radiation, \(\theta \approx \gamma^{-1}\), optimized for the first mode, \(r=1\), then we have for the \(r\)th resonance

\[
F_2 = 4 \sin^2 \left(\frac{\pi(1 + \gamma^2 \theta_r)}{4}\right). \tag{5}
\]

Note that working far below the Cherenkov radiation, i.e., if either \(\varepsilon < 1\) or \(\varepsilon - 1 \ll \gamma^2\), which is always the case in our situation, the condition of the \(r\)th resonance [derived from Eq. (1)],

\[
\theta_r + \gamma^2 = \frac{2r\lambda}{l}, \quad l = \lambda \gamma^2, \tag{6}
\]

gives

\[
\frac{\theta_r}{\gamma^2} = \frac{(2r-1)}{\gamma^2} = \theta_1^0(2r-1), \tag{7}
\]

so that \(F_2 = 4 \sin^2(r \pi/4)\), i.e.,

\[
F_2 = 4 \quad \text{for } r = 1, 3, \ldots \text{ and } F_2 = 0 \quad \text{for } r = 0, 2, 4, \ldots . \tag{8}
\]

Thus, the even modes \(r\) are suppressed; be reminded that this is due to fact that at each neighboring interface the TR waves have different signs, since \(\Delta \varepsilon\) changes its sign at each successive interface.

Finally, for a single electron traversing \(M\) periods, each period consisting of two adjacent layers of alternating lossless materials, the multilayer factor \(F_3\) for coherent summation of TR from each period is

\[
F_3 = \frac{\sin^2(MX)}{\sin^2(X)}, \tag{9}
\]

where \(X\) is a half-phase difference between each period; in our case

\[
X \approx \frac{l \pi(\gamma^2 + \theta_r^2)}{2\lambda} \approx \frac{\pi(1 + \gamma^2 \theta_r^2)}{2}. \tag{10}
\]

The factor \(F_3\) exhibits the resonances at the modes \(r\) occurring at \(X_r = r \pi\). Since the number of periods should be high, \(M \gg 1\), the radiation pattern of the multilayer structure will be concentrated very nearly the \(r\)th resonant angle.
\[ \theta_r = \gamma^{-1} \sqrt{2r - 1}, \]  

so that in the close vicinity of each resonance, \( \theta = \theta_r + \Delta \theta \), with \( \Delta \theta \ll \theta_r \ll 1 \), we can write

\[ X_r = r \pi + \pi \gamma \sqrt{2r - 1} \Delta \theta \]  

and

\[ (F_3)_r = \frac{\sin^2(M \pi \gamma \sqrt{2r - 1} \Delta \theta)}{(\pi \gamma \sqrt{2r - 1} \Delta \theta)^2}, \quad r = 1, 3, 5, \ldots. \]  

Notice that in the case of exact resonance, \( X = 0 \), or \( \Delta \theta = 0 \), we can write (9) and (13) in the form

\[ F_3 = M^2. \]  

According to Eq. (4) we can write now

\[ \frac{dN_r}{d\Omega d\omega} = \frac{\alpha(\Delta e)^2 \gamma^4 (2r - 1) \sin^2(M \pi \gamma \sqrt{2r - 1} \Delta \theta)}{4r^4 \pi^2 \alpha \pi \gamma \sqrt{2r - 1} \Delta \theta^2}. \]  

The total number of the photons in all the modes at the frequency \( \omega \) is

\[ N_\omega = \sum_{r=1}^{\infty} N_r \quad \text{with} \quad r = 2k - 1. \]  

From (16) we have

\[ \frac{dN_r}{d\omega} = \frac{\alpha(\Delta e)^2}{2 \pi \omega} M^4 \gamma^4 \]  

which shows that the main mode with \( r = 1 \) carries \( \sim 92\% \) of the entire radiation, so the rest of the modes are insignificant. Anyway, summing up all the modes up to \( r = 101 \), we find

\[ \frac{dN_\omega}{d\omega} \approx 1.1M^4 \gamma^4 \frac{\alpha(\Delta e)^2}{2 \pi \omega}. \]  

Equation (20) can be modified to take the photoabsorption into account as follows:

\[ \frac{dN_\omega(\omega)}{d\omega} = 1.1 \gamma^4 \frac{\alpha(\Delta e)^2}{2 \pi \omega} L_{\text{att}}(1 - e^{-\beta \omega / L_{\text{att}}}), \]  

where \( L_{\text{att}} = 2(L_1^{-1} + L_2^{-1})^{-1} \) is the mean attenuation length, with \( L_i \) being the attenuation lengths of the materials comprising the structure.5

As we show below, to measure the TR one should use spatial filtering. Thus, the experimental data would be better characterized not by the radiation efficiency integrated over all modes and all angles but rather by TR angular distribution in the vicinity of the first resonance. In this case, the TR differential efficiency with photoabsorption and electron scattering taken into account can be calculated using Eqs. (2)–(5) where instead of Eq. (13) for \( F_3 \) one should use \( \tilde{F}_3 \)

\[ \tilde{F}_3 = e^{-(M-1)(\xi + \rho)} \cos h[M(\xi - \rho) / \cos h(\xi - \rho) / \cos h(2X)], \]

where \( \xi = (\mu_i l_i + \mu_f l_f) / (2 \cos \theta) \) is the mean photoabsorption coefficient of the structure, \( \mu_i \) is the photoabsorption coefficient of the \( i \)th material, \( i = 1, 2, \mu = \pi / (2L_\omega) \), \( L_\omega = \int (dE/dz)(l_1(l_2 l_3) + (dE/dz)(l_2 l_3)) \) is a distance at which nearly all electrons become scattered or absorbed, and \( dE/dz \) are the electron-atom collision losses per unit length calculated as

\[ \frac{dE}{dz} = \frac{2\pi e^2}{\beta^2} \gamma Z_{\text{Ai}} \left[ \ln \left( \frac{(\gamma - 1)^2 (\gamma + 1)}{2l_i^2} \right) + (\gamma - 1)^2 (\gamma - 1) \ln 2 + \frac{1}{\gamma^2} \right], \]

where \( m \) is the rest mass of the electron, \( r_e = e^2/(mc^2) = 2.82 \times 10^{-13} \) cm is the classical electron radius, \( e \) is the electron charge, \( l_i = 9.73Z_i + 58.8Z_i^{-0.19} \) eV is the approximate ionization potential of the atom (for atomic number \( Z_i > 13 \)), and \( N_{\text{Ai}} \) are the atomic number densities.

**IV. DESIGN OF EXPERIMENT**

Aiming to design a proof-of-principle experiment, we have chosen as a radiator an element widely used in the x-ray technology (e.g., for x-ray and EUV mirrors), whose properties are well known and which is readily available, molybdenum \((Z=42)\), and which has \( K \) shell at \( E_K = 19.9995 \) keV. Next, we conducted a search of the spacer, which revealed a number of materials suitable for the Mo radiator. These materials are Pb, Ag, Tl, and Th, which provide a very convenient set of options; for instance, lead (Pb) as a spacer has a great advantage of being accessible and technologically easily manageable. Another advantage also is that lead has a great tolerance to mismatching of the crystal lattice constants (this mismatch usually creates problems with mechanical stability of multilayer structures). The contrast of dielectric constants, \( |\Delta \epsilon(\omega)|^2 \) for Mo–Pb, Mo–Ag, Mo–Tl, and Mo–Th pairs, shown in Fig. 1 demonstrates a well pronounced peak near \( E_K \) of Mo, and the wing-matching condition is clearly satisfied, which differs significantly from the light spacer + heavy radiator structures (e.g., Mo–Si, upper curve in Fig. 1).

Let us focus on the E-beam of 30 MeV energy and calculate the transition radiation efficiency, integrated over all the angles and modes, using Eq. (21), which takes the photoabsorption into account. Note that the effects of electron scattering are negligible as compared to the photoabsorption for the TR structures and E-beams under study [\( L_\omega \gg L_{\text{att}} \), see Eqs. (21) and (23)].

The TR spectra (Fig. 2) for the most efficient couples, Mo–Ag, Mo–Tl, and Mo–Pb, clearly demonstrate the
strongly pronounced resonant line, with contrast between the wings and the center easily reaching two orders of magnitude. The radiation lines shown in Fig. 2 have ~1% total linewidth. One can see that even though the peak value of \( \mu Mo-Pb \) is largest for Mo–Tl structure (Fig. 1), with photoabsorption taken into consideration the most efficient couple of radiator and spacer is Mo–Ag (Fig. 2), which we focus at in the rest of our paper.

As the thickness of structure increases up to the attenuation length and beyond it, the photon output is getting saturated. In practical terms, it would be reasonable to keep up to ~1000 periods in Mo–Ag structure, as one can see in Fig. 3 that depicts the total number of generated TR photons within a resonant line versus the total thickness of the structure. The total number of photons per electron in the Mo–Ag nanostructure is about \( 3 \times 10^{-7} \).

It is easy to see that this efficiency is high enough for medical applications discussed in Sec. I. Indeed, using this structure with, e.g., a reconditioned medical accelerator Clinac (see, e.g., Ref. 18) at 30 MeV electron energy, 120 Hz repetition rate, and 2 \( \mu s \) pulse duration, we arrive at \( 6 \times 10^7 \) photons/s flux. This flux is essentially the same as the one utilized in the prototype bichromatic x-ray contrast diagnostics (see, e.g., Ref. 5); however, instead of a large synchrotron used in that prototype, TR from a Mo–Ag nanostructure involves a modest-size medical electron accelerator, affordable to any large hospital.

V. BREMSSTRAHLUNG AS A NOISE

At high electron and photon energies there is another major radiation process competing with TR: it is bremsstrahlung.\(^3\) In the soft x-ray domain bremsstrahlung is significantly weaker than TR,\(^5,6\) whereas for hard x rays it should be taken into account. As we show below, spatial filtering must be used for efficient detection of TR generated by 30 MeV electrons traversing a Mo–Ag nanostructure. In this section we study the angular distribution of both TR and BS. BS differential efficiency can be calculated in the Born approximation with screening taken into account using, e.g., the Thomas-Fermi model.\(^3\) Including the photoabsorption into consideration, bremsstrahlung efficiency can be calculated as

\[
\frac{dN_{bs}}{d\Omega d\omega} = \frac{dN_{s}}{d\Omega d\omega} \left[ \frac{1 - \exp(-\mu_{Ag}L)}{\mu_{Ag}L} + \frac{dN_{s}}{d\Omega d\omega} \left[ \frac{1 - \exp(-\mu_{Mo}L)}{\mu_{Mo}L} \right] \exp(-\mu_{Ag}L) \right],
\]

where

\[
\frac{dN_{s}}{d\Omega d\omega} = 4\pi Z^2 \gamma^2 N_0 L \left( \frac{4}{\pi x^2 + 1} \right)
\times \left[ (x^4 + 1) \log \left( \frac{111(x^2 + 1)}{Z^{1/3} - (x^2 - 1)^2} \right) \right],
\]

\( \gamma = E_0/mc^2, \ E_0 = 30 \text{ MeV}, \ \mu \) is the photoabsorption coefficient, \( x = \gamma \theta, \ N_0 = \) the number density of material, and \( L = lM/2 \) is the half-width of the whole nanostructure. In Eq. (24) \( dN_{s}/d\Omega d\omega \) should be calculated separately for Ag and Mo using Eq. (25). Figure 4 shows differential efficiency of TR and BS for 750-period Mo–Ag nanostructures in the photon energy range around the \( K \)-shell photoabsorption edge of Mo. One can see that the peak of TR radiation efficiency is about four times higher than the efficiency of BS radiation.

VI. DETECTORS AND MEASUREMENT ARRANGEMENT

In practice, the TR can be detected through a pinhole (see, e.g., Ref. 6), opening a small solid angle around the TR peak. Figure 5 shows angular distribution of differential efficiency of BS and TR at the TR peak energy \( \approx 20 \text{ keV} \) for 750- and 250-period Mo–Ag structures. Let us consider the 250-period nanostructure and calculate the diameter of a circular pinhole suitable for experimental observation of TR. The diameter, corresponding to the half-width of the TR radiation peak would be \( d = R \delta \theta_{TR} \), where \( \delta \theta_{TR} \approx 6.1 \times 10^{-3} \) rad (see Fig. 5) and \( R \) is the distance from the Mo–Ag structure to the pinhole. For instance, when \( R = 200 \text{ cm}, \) the pinhole diameter \( d \approx 0.012 \text{ cm} \).

Since bremsstrahlung radiation is quite strong in the hard x ray and its bandwidth is much broader than that of TR, the...
Choice of x-ray detector also requires special attention. To estimate the level of BS noise, let us calculate the number of BS photons passing through the pinhole registered by the detector. The most suitable for energy range under the study are Si and CdTe detectors. We chose Amptek XR-100CR-Si detector. The most suitable for energy range under the study is XR-100T-CdTe-1 mm detectors and calculate the number of registered BS photons in accordance with the detector efficiencies using the following scheme. Below 5 MeV the detection efficiency $E$ of detectors is known, so that the number of detected photons is

$$\frac{dN_{\text{det}}}{d\Omega} = \int_{E_1}^{E_2} \frac{dN_{\text{br}}}{d\omega} E(\omega) d\omega.$$  \hfill (26)

From 5 to 30 MeV the main mechanism contributing to the detection events is pair production:

$$\frac{dN_{\text{det}}}{d\Omega} = N_0 d_D \int_{E_1}^{E_2} \frac{dN_{\text{br}}}{d\omega} \sigma_{\text{pair}}(\omega) d\omega,$$  \hfill (27)

where

$$\sigma_{\text{pair}}(\omega) = r^2 Z^2 \alpha \left[ \frac{28}{9} \log \frac{2h\omega}{mc^2} - \frac{218}{27} \right].$$  \hfill (28)

d$_D$ is the detector thickness (500 $\mu$m and 1 mm for Si and CdTe detectors, respectively), and $N_0$ and Z are number density and atomic number of detector material (Si or CdTe). The total number of registered BS photons is the sum of the two contributions given by Eqs. (26) and (27). For the pinhole of the diameter $d=0.012$ cm (as discussed above) placed at the angle of TR maximum ($\theta_{\text{TR}}=0.9457^\circ$) the detectors would register the following numbers of BS photons per electron passing through the pinhole: $1.62 \times 10^{-9}$ (XR-100CR-Si) and $7.89 \times 10^{-9}$ (XR-100T-CdTe). Thus, competing with the TR bremsstrahlung can be reduced sufficiently enough to detect the TR signal with a good margin.

VII. COHERENCY AND PRECISION TOLERANCE OF MULTILAYER STRUCTURE

Unavoidable defects and manufacturing inaccuracies (e.g., width fluctuation of the layers) in periodic structures lead to inhibition of coherent summation of TR from each period. Intensity of a signal resulted from ideally coherent summation at the exact resonant angle, $X=0$, is proportional to $M^2$ [see Eq. (14)], whereas in the opposite case of the fully incoherent structure the expected intensity is $\sim M$ [similar to the integration over all the modes; see Eq. (20)]. The theory of randomization-induced inhibition of coherency of multielement arrays, whereby the spacings between adjacent elements randomly deviated from the desired periodicity $l_0$ has been developed in Ref. 22. It has been shown that regardless of the specific statistical distribution of random width deviation in the structure, the resulting coherency is described with high accuracy by Gaussian distribution. The total gain, or the factor $M^2$ in Eq. (14), should now be replaced by

$$E_M M,$$  \hfill (29)

where $1 \leq E_M \leq M$ with $E_M$ being a coherent enhancement.

Following Ref. 22, we introduce a normalized standard deviation $\sigma_0$ of the spacing from the designed spacing $l_0$ between two adjacent periods of the structure with

$$\sigma_0 = \frac{\sigma_i}{l_0},$$  \hfill (30)

where $\sigma_i$ is a standard deviation, and define the randomiza-
tion parameter $r$ and the inverse to it, coherency range, $M_{coh}$ as
\[ r = \frac{1}{\pi \sigma_0^2} \] and $M_{coh} = \frac{1}{r}.$
(31)

At the angle of exact resonance, $X=0$, the following relationship is valid:22
\[ \mathcal{E}_M(X=0) = \frac{(M/2) \sinh(2r) - e^{-rM} \sinh(rM)}{M \sinh(r)} \]
(32)

The behavior of $\mathcal{E}_M$ as function of $M$ for various standard deviations $\sigma_0$ is depicted in Fig. 6. Assuming at least some coherence, i.e., $r \ll 1$ or $M_{coh} \gg 1$, and the exact solution (32) can be reduced to a simpler formula:
\[ \mathcal{E}_M(X=0) = M_{coh} \left[ 1 - \frac{e^{-r} \sinh \zeta}{\zeta} \right], \quad \zeta = \frac{M}{M_{coh}}, \]
(33)
which indicates that even when the ideal coherence is lost because of $M \gg M_{coh}$, we still have $\mathcal{E}_M \approx M_{coh} \gg 1$, i.e., a substantial enhancement over the case of fully random structure. Equation (33) shows that when $\zeta = 1$ or $M = M_{coh}$ the system exhibits only $\sim 50\%$ loss of coherency compared to the ideal case, thus still providing for the orders of magnitude enhancement over the fully randomized case.

It was also shown in Ref. 22 that both (32) and (33) can be approximated within better than 5% by much simpler heuristic formula, which is intuitively transparent and could be useful for practical applications in the entire range of parameters $M$ and $M_{coh}$:
\[ \mathcal{E}_M \approx \frac{MM_{coh}}{M + M_{coh} - 1} = \frac{M}{(M-1)r + 1}. \]
(34)

Let us calculate the coherency enhancement and coherency length for the Mo-based nanostructures studied above. For 10 MeV electron beam we have $l \approx 27$ nm. Assuming \( \sigma_l = 2.7 \text{ Å} \) tolerance in the period thickness, one has the standard relative deviation $\sigma_0 = 1\%$ and the coherency range $M_{coh} \approx 10^3$, so with the total number of periods $M \sim 10^2$, one has a gain, $\mathcal{E}_M$, only 7% below the ideally coherent gain $M^2 = 10^4$. With $M \sim 10^3$, one has the total gain $\sim 50\%$ of the fully coherent $M^2 = 10^6$. With the electron energy of 30 MeV, $M = 750$, and for $\sigma_l = 2.23$ nm and $\sigma_0 = 6.69$ nm one has about 60% and 14% of the fully coherent gain, respectively. Using Eq. (34) one can also calculate manufacturing precision tolerance for the multilayer nanostructure period allowing to have specific coherency gain. For example, for the 750-period nanostructure one should have $\sigma_l \leq 1$ nm in order to keep the TR intensity within 10% of its maximum (fully coherent) signal.

VIII. CONCLUSION

We have proposed to use the resonant transition radiation of metal-metal multilayer nanostructures in the vicinity of photoabsorption edge of constituent materials as a narrow-line source of hard x rays. Detailed analysis revealed that for hard x rays the optimal spacer-radiator pair of materials comprising the nanostructure should have the spacer, as a rule, with its atomic number larger than those of the radiator in order to eliminate the dielectric constant contrast at the far wings of the radiation line. We have applied the proposed criterion to select candidates on creation of efficient hard x-ray radiation source. For a model pair, Mo (radiator) and Ag (spacer), we have calculated the TR radiation efficiency, which shows a narrow-line near the photoabsorption edge of Mo of about 1% width. One of the most important factors in attaining an efficient source of resonant transition radiation based on atomic absorption edges becomes filtering out the Bremsstrahlung radiation. We demonstrate that for Mo–Ag couple with E-beam of 30 MeV the TR line can be well resolved when both spatial and spectral filtering is used.

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3W. Sharf, Biomedical Particle Accelerators (APS, New York, 1994), Chap. 6.
10The physics of resonant anomalous x-ray dispersion and scattering (with the index of refraction being part of it) has been developed in great detail in the context of crystallography, starting from A. H. Compton and S. K. Allison, X-ray in Theory and Experiment (Van Nostrand, New York, 1935); the excellent recent collection of articles can be found in Resonant Anomalous X-ray Scattering: Theory and Application, edited by G. Materlik, C. J. Sparks, and K. Fisher (North-Holland, New York, 1994). In particular, the detailed review of experimental data and earlier work are found in the papers by B. Lengeler ibid., p. 35; R. L. Blake, J. C. Davis, D. E. Graessle, T. H. Burbine, and E. M. Gullikson, ibid., p. 79; and D. H. Templeton, ibid., p. 1. See also http://www-cxro.lbl.gov/optical_constants/
13M. L. Ter-Mikaelian, High Energy Electromagnetic Processes in Con-
For our calculations we have chosen the E-beam of 30 MeV because one of the best candidates for our future proof-of-principle experiment is Linac at Brookhaven Accelerator Test Facility, which has operating energy above 25 MeV. It must be noted, however, that for many reasons (design, accelerator accessibility, technology, bremsstrahlung noise, etc.), it is actually preferable to have lower E-beam energies, near 10 MeV.