InAsSb-Based Infrared Photodetectors: Thirty Years Later On

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Received: 24 September 2020; Accepted: 9 November 2020; Published: 9 December 2020

Abstract: In 1989, one author of this paper (A.R.) published the very first review paper on InAsSb infrared detectors. During the last thirty years, many scientific breakthroughs and technological advances for InAsSb-based photodetectors have been made. Progress in advanced epitaxial methods contributed considerably to the InAsSb improvement. Current efforts are directed towards the photodetector’s cut-off wavelength extension beyond lattice-available and lattice-strained binary substrates. It is suspected that further improvement of metamorphic buffers for epitaxial layers will lead to lower-cost InAsSb-based focal plane arrays on large-area alternative substrates like GaAs and silicon. Most photodetector reports in the last decade are devoted to the heterostructure and barrier architectures operating in high operating temperature conditions. In the paper, at first InAsSb growth methods are briefly described. Next, the fundamental material properties are reviewed, stressing electrical and optical aspects limiting the photodetector performance. The last part of the paper highlights new ideas in design of InAsSb-based bulk and superlattice infrared detectors and focal plane arrays. Their performance is compared with the state-of-the-art infrared detector technologies.

Keywords: infrared detectors; InAsSb; superlattice; higher operating temperature

1. Introduction

The development of InAs1−xSbx (InAsSb) has a long history. InAsSb was first synthesized in 1958 by Woolley and Smith [1]. This III–V material belongs to the family of narrow gap semiconductor compounds. Among them we can also distinguish IV–VI (Pb1−xSnxTe), and II–VI (Hg1−xCdxTe) infrared (IR) material systems. The bandgap of these semiconductors and hence the spectral response of the devices can be tuned for specific detectors’ applications. The favorable properties of narrow-gap semiconductors: high electron mobility, high optical absorption coefficient, and low thermal generation rate together with the capability for bandgap engineering, make these compounds almost perfect and ideal material for the design of detectors that are applied in a wide range of IR spectrum.

The most important position in IR detector technology is held by HgCdTe discovered in 1959 in the UK by Lawson, Neilsen, Putley, and Young [2]. HgCdTe has triggered the development an unprecedented degree of freedom in IR detector design and has inspired the development of the three “generations” of detectors considered mainly for military applications [3].

The first discoverers of HgCdTe also knew [4], and the explosions they experienced would have confirmed it, that the Hg vapor pressure over HgCdTe, which can be much greater than 20 atmospheres, would be a severe problem for crystal growth. The need to provide high mercury vapor pressure...
has been a major drawback during the growth of HgCdTe and for this reason alternative detector technologies were attempted over the past sixty years. One of those was PbSnTe, which was thoroughly studied in parallel with HgCdTe in the late 1960s and early 1970s [5–7]. Good quality long-wave infrared (LWIR) photodiodes were demonstrated as PbSnTe was comparatively easy to grow. However, in the late 1970s there were two reasons which contributed to the termination of the research on PbSnTe: large temperature coefficient of expansion (TCE) mismatch with silicon and high dielectric constant. Relatively fast response time was required in scanned IR imaging systems of the 70s last century so that the scanned image is not blurry in the scan direction.

Nowadays with the trend towards focal-plane arrays (FPAs) this drawback does not play a key role like during the very first-generation systems’ design. In turn, due to large TCE, the indium bonds between silicon readout and the detector array in hybrid structure are exposed to adverse stresses during repeated cooling cycles from room temperature to cryogenic temperature. Today this drawback is omitted by fabrication of monolithic PbS and PbSe polycrystalline arrays on Si substrates.

The report on InAs$_{0.8}$Sb$_{0.15}$ photodiodes grown by step-graded liquid phase epitaxy (LPE) technique [8] initiated extensive research on InAsSb IR photodetectors in the mid 1970s. Near-lattice-matched InAs$_{1−x}$Sb$_x$ (0.09 $≤$ x $≤$ 0.15) device structures grown on GaSb substrates revealed the best parameters of InAs$_{1−x}$Sb$_x$ photodiodes [9]. InAsSb has the smallest bandgap of conventional III–V semiconductors and can be used as active material for mid-wavelength infrared (MWIR) and LWIR photodetectors. However, the energy gap of InAs$_{1−x}$Sb$_x$ during the 1970s was not exactly controlled in the middle composition range. In consequence, the InAsSb did not have a sufficiently small gap at lower temperatures, especially 77 K, for operation in the 8–14 $\mu$m wavelength range. Newly designed III–V strained-layer superlattices (SLs) consisting of a number of alternating thin mismatched crystal layers were developed to face this disadvantage.

Low dimensional solids representing new generation of materials for IR detectors have been proposed since the 1970s. Only a few years after the first GaAs/AlGaAs quantum heterostructures, the HgTe/CdTe SLs system was demonstrated in 1979 [10]. Two additional structures were subsequently introduced: InSb/InAsSb SLs with strain-induced bandgap reduction [11] and InAs/GaInSb SLs with superlattice-induced band inversion [12–14]. Even smaller bandgaps can be obtained than either constituent if InAs$_{1−x}$Sb$_x$ is combined with InSb or InAs in a SL. The first high-detectivity InAs$_{0.15}$Sb$_{0.85}$/InSb SLs photodiodes with detectivity $≥1 \times 10^{10}$ cmHz$^{1/2}$/W at wavelength, $\lambda = 10$ $\mu$m and 77 K were fabricated in 1990 by Kurtz et al. [15]. InSb/InAsSb SLs grown on InSb substrates revealed the excessive net stress, which inevitably caused cracks of the epitaxial material, making the fabrication of detector arrays impossible at that time [16].

The next step in development of SLs structures for IR detectors was concentrated on InAs/GaInSb material [12,13]. The larger valence band energy differences between InAs and GaSb than that between InSb and InAsSb directed mainstream research towards InAs/GaInSb type-II superlattices (T2SLs) allowing larger absorption in LWIR spectral range.

Historically, the InAs/InAsSb SLs as material for IR detectors had been selected more than 20 years after InSb/InAsSb ones [17]. Shorter Shockley-Read-Hall (SRH) minority carrier lifetimes of InAs/GaInSb T2SLs, were the main motivation to revive research into InAs/InAsSb T2SLs [18]. In addition, InAsSb-based detectors could be used in many civilian and military applications. The topic of InAsSb-based IR detectors has been covered only in two review papers [19,20] and one book [21]. In the last year, Steenbergen published an excellent chapter with the main emphasis on growth methods, material properties, and device fabrications [20]. In the present paper more attention is directed on the position of InAsSb-based detectors in confrontation with the present stage of IR detectors offered on global market. This evaluation is supported by theoretical estimates and experimental data.

2. Brief View on Crystal Growth of InAsSb-Based Materials

The InAs$_{1−x}$Sb$_x$ properties were first investigated by Woolley and coworkers in the late 1950s. They determined InAs-InSb pseudobinary phase diagram [1], the miscibility [1], and identified...
the variation of properties such as bandgap \[22,23\] and effective masses on composition \[24–26\]. Polycrystalline InAsSb samples were measured to determine fundamental parameters. Many problems related to the InAsSb crystal growth arise from the pseudobinary phase diagram shown in Figure 1. The very low diffusion rates in the solid phase, the large separation between the liquidus and solidus, and the lattice mismatch (6.9% between InAs and InSb \[a_o(InAs) = 0.60584 \text{ nm}, a_o(InSb) = 0.64794 \text{ nm}\]) are challenging for the crystal growth.

![Figure 1. Pseudobinary phase diagram for the InAs-InSb systems [1].](image)

Progress in InAsSb ternary system has been limited by crystal synthesis problems. These difficulties are being overcome systematically by epitaxial methods such as: LPE, molecular beam epitaxy (MBE), and metalorganic chemical vapor deposition (MOCVD). Bulk single crystals are prepared mainly for compositions close to the binary compounds. A wide range of topics are gathered in thorough studies on electronic and photonic materials published in the Springer Handbook of Electronic and Photonic Materials \[27\].

LPE is a relatively simple high-quality technique, with less expensive epitaxial equipment and high utilization rate of the source material. It is particularly suitable for the preparation of thick-film layers. Since LPE technique is a near-thermodynamic equilibrium growth method, it cannot be used for the growth of the metastable ternary and quaternary antimonide compounds with miscibility gaps like InAsSb. The growth rate of LPE varies typically of 100 nm/min to a few \mu m/min (depending on substrate crystalline phases) and is generally higher than MBE and MOCVD. However, the controlled growth precision of very thin epilayers, especially quantum-well (QW) devices, SLs, and other complex microstructure materials cannot be implemented by LPE. In addition, the surface morphology of MBE or MOCVD grown layers is usually better than that grown by LPE.

The era of MBE and MOCVD growth of III–V semiconductors began in the early 1970s. It is often difficult to decide which epitaxial growth technique, MBE or MOCVD, to choose. Each of them has specific advantages in a given device application. Table 1 presents some characteristics related to the different classes of methods for III–V compounds \[28\].

| MBE | MOCVD |
|-----|-------|
| Long individual campaigns, no setup variability. | Longer individual campaigns, less setup variability. |
| Uniformity easier to tune, largely set by reactor geometry. | More flexibility for source and reactor geometry. |
| Excellent quality/crystallinity. | Uniform or MOCVD grown layers is usually better than that grown by LPE. |
| High utilization rate of the source material. | Shorter maintenance periods. |
| Growth near thermodynamic equilibrium, no hydrogen passivation, no burn in inherent to fusion cells toward the substrate in the form of streams being chemically unchanged. | No hydrogen passivation, no burn in inherent to fusion cells toward the substrate in the form of streams being chemically unchanged. |
| The use of carrier gas is not necessary-the inherently long mean free paths result in highly directional elemental beams and the UHV environment ensures high material purity. Fast shutters enable deposition of more sophisticated layer structures like SLs and QWs. The valved sources are generally used because of relatively high vapor pressures for V group elements. Gas sources (e.g., CBr\(_4\) for C doping), precursor sources (e.g., GaTe for Te doping), and plasma sources (e.g., nitrogen plasma) are applied. |
Table 1. MOCVD and MBE methods comparison (after [28]).

| Category       | MBE                                                                 | MOCVD                                                                 |
|----------------|----------------------------------------------------------------------|------------------------------------------------------------------------|
| **Growth**     | Fast switching for proper interfaces.                                | High growth rate for bulk layers.                                      |
|                | Able to grow thermodynamically forbidden materials.                  | Growth near thermodynamic equilibrium, excellent quality/crystallinity.|
|                | No hydrogen passivation, no burn in inherent to MOCVD.              | Ability to control background doping.                                   |
|                | Uniformity easier to tune, largely set by reactor geometry.          |                                                                        |
| **Economic**   | Longer individual campaigns, less setup variability.                | Shorter maintenance periods.                                           |
|                | Overhead does not scale with run rate. Contribution per wafer Increases with wafer volume. Economic to idle. Overhead cost scales with run rate. |
|                | Lower material cost/wafer.                                          | Higher safety risk, increasing scrutiny of legislative bodies worldwide.|

MBE enables the control of the composition of the growing crystal with monolayer (ML) resolution due to the both slow growth rate (~1.0 ML per second) and mounting the shutters just in front of the crucibles. Calibration of the growing crystal with the electron beams is possible because of low background pressure in MBE. The layer-by-layer growth mode, III/V flux ratio, as well as the quality of the growing substrate can be determined by the reflection high energy electron diffraction (RHEED) technique. MBE provides the ability to control the interfaces by shutter sequence with a precision of 0.1 s because the growth proceeds in thermodynamic imbalance conditions. The leaks can be a major problem because the growth proceeds under extremely low pressure. The growth chamber is predominantly cooled by liquid nitrogen to strengthen high vacuum by preventing molecules from peeling off from the chamber walls.

MOCVD is another important growth technique widely used for heteroepitaxy of the QW structures and SL. Similarly to MBE, it also enables the growth of heterostructures with abrupt interfaces between the individual layers. The growth in MOCVD technique proceeds on a heated substrate but in a much higher pressure than MBE (typically 15 to 750 Torr). There are several types of MOCVD reactor designs. The growth proceeds at near atmospheric pressure in the atmospheric MOCVD and this requires the use of a large amount of carrier gases. In turn, the reactor pressure is kept low in the low pressure MOCVD and in this case the growth rate is then slower as in the MBE. Strict security precautions must be incorporated in the MOCVD laboratory. In order to avoid any deadly accidents, many safety precautions have to be implemented because the precursor gases and the carrier gases are often highly toxic or explosive. In almost all semiconductor fabrications, safety and environmental concerns are important issues because the growth processes are almost always associated with toxic and dangerous materials.

MOCVD growth proceeds not by physical deposition but by chemical reaction. In contrast to MBE, MOCVD technique requires more complex compound precursors, namely metal-organic sources (e.g., di-methyl or tri-ethyl Cd, Te, Al, etc.), hydrides (e.g., PH3, etc.), and other gas sources (e.g., Si2H6). The partial pressures of the precursors are controlled with mass flow controllers. In MOCVD, the precursors are transported over the substrate where they pyrolyze resulting in epitaxial growth. MOCVD requires the use of a carrier gas (typically hydrogen) to transport the precursors across the wafer surface. The valve actuation for varying injection ports of a gas manifold enables the growth of
complex heterostructures. The fabrication of optoelectronics devices involving thermodynamically metastable alloys is dominated by both MBE and MOCVD techniques.

From the economical point of view (see Table 1 and [28]):

- the cost characteristics are very different because of the specific requirements:
  - for MBE the overhead is relatively fixed and does not vary with volume,
  - for MOCVD, the overhead costs tend to vary with production volume, and therefore
- MOCVD exceeds in a case of significant overcapacity (long idle time), and the opposite is true for MBE which wins on a cost basis when fully loaded,
- the two methods are very similar considering the production efficiency time,
  - for MBE, the idle time lasts on average several months where the reactor is down,
  - MOCVD service procedures are much more frequent but less time consuming, and therefore
- for MBE system, significant bake times are needed when the growth chamber must be opened for the service repairs,
- in contrast, MOCVD does not require long bake times (MOCVD is able to recover more quickly from failures).

Manasevit and Simpson firstly performed epitaxial growth of antimonides thin layers using MOCVD with TMGa (trimethylgallium) and SbH$_3$ (stibine) precursors for GaSb films deposition in 1969 [29]. At present, the typically used III-group metal-organic sources by MOCVD for antimonide-based materials are 3-methyl compound and 3-ethyl compound, such as: TMAI, TMIn, TMGa, TEIn, TEGa, etc. [30,31]. In turn, V-group commonly used precursors are: AsH$_3$, PH$_3$, TMBi, TMSb, and RF-N$_2$, etc. The growth temperature of antimonides (low melting point materials) is close to 500 $^\circ$C. It appears that below 500 $^\circ$C the vast majority of III-group metal-organic precursors cannot decompose in 100%. Therefore, new organic source materials with a lower decomposition temperature are introduced including: TDMASb (trisdimethylaminoantimony), TASb (triallyantimony), TMAA (trimethylamine alane), TTBAI (tritertiarybutylaluminum), EDMAA (ethyldimethylaminealane), etc. In the case of Al-containing antimonide materials, carbon and oxygen contamination problems exist. The active hydrogen atoms’ absence on the surface of epitaxial layers is expected to be the reason of this effect. Carbon is typically p-type doping impurity, which causes certain difficulties in growing of n-type doping Al-containing epitaxial layers.

MBE grown antimonides was first reported in the late 1970s [32,33]. In comparison with GaAs and other arsenides, the growth of GaSb-based is characterized by relatively low Sb vapor pressures, or, equivalently, by its high sublimation energy. GaSb and AlSb are nearly lattice-matched mutually and to InAs and for this reason they are the subject of intensive research. The substrate temperature during the growth of GaSb and AlSb is usually between 550 and 600 $^\circ$C. MBE reduces the concentration of O-doping and avoids the C-contamination issue in Al-based materials growing by MOCVD. Most of low-dimensional structures (quantum wires-QRs, QWs, and quantum dots-QDs) and devices having complex structures were first grown by MBE. It was experimentally proven that the implementation of crystallographically misoriented substrates (small angle offset) contributed to higher-quality epitaxial layers [34].

The growth of antimonide-based III–V epitaxial layers is usually performed on InSb, InAs, and GaSb low-defect substrates. GaAs and GaAs-coated Si substrates and other heterogeneous substrate materials for epitaxy were introduced to face the problem that antimonides have no semi-insulating substrate. The growth of multilayers with abrupt but incremental compositional shifts between adjacent layers and continuous compositional grading of thick epilayers enables one to obtain a variety of substrate structures. Wafer bonding techniques and selective removal of the seeding substrate methods can be combined in IR detector fabrications. With high-quality substrates with
increased functionality and bandgaps and lattice constants differing significantly more than feasible with binary compound wafers (e.g., GaSb or InAs), it seems possible to implement [35].

The initial efforts in fabrication and characterization of InAsSb material until 1994 were described in two Rogalski papers [19,21]. The book chapter written recently by Steenbergen [20] has provided updated information on the growth methods and characterization techniques of InAsSb bulk and SLs materials for the last thirty years. Because of this, here only main trends in material fabrication are covered.

Current interest is directed toward the InAsSb photodetectors cut-off wavelength (\(\lambda_{\text{cut-off}}\)) extension beyond that available when lattice-matched or lattice-strained to binary substrates; mainly GaSb. In order to improve FPAs manufacturability and increase their size, low-cost GaAs and Si substrates are used. The Si substrates are very convenient in IR FPA technology because the coupling with Si readout enables fabrication of very large arrays exhibiting the prolonged thermal cycle reliability (Si substrates are cheaper and available in large area).

In practical design of InAsSb-based device structures, a serious problem is the control of photodetector’s \(\lambda_{\text{cut-off}}\). The accuracy of the bandgap depends on material composition, stress in the buffer layer between substrate and the active region, the crystalline quality of the ternary samples, (potentially CuPt ordering) which is determined by the growth method and conditions, and finally the bandgap measurement uncertainty. Different techniques used in bandgap characterization (absorption, ellipsometry, photoluminescence (PL) and electroluminescence (EL)) and experimental results are given by Steenbergen [20].

Usually, the spontaneous ordering of CuPt in mid-composition range of InAs\(_{1-x}\)Sb\(_x\) causes a problem. Better alloy composition uniformity of group V materials on lattice-mismatched substrates is required to grow controlled, high crystalline quality InAsSb without spontaneous ordering [36]. In the technology of ternary semiconductors, atoms of group V having different sticking coefficients are more difficult to control than group III ternaries [37,38]. Using modulated MBE (MMBE) technique with arsenic and antimony shutter modulation (see Figure 2), better control of InAsSb composition is reached.

![Figure 2](image_url) The scheme of the MMBE shutter sequence during the growth of InAs\(_{0.76}\)Sb\(_{0.24}\) and the surrounding InAs layers. The filled bars denote intervals when the corresponding shutters are open (after [37]).

The spontaneous CuPt ordering during InAsSb MBE growth is probably caused by residual strains [39]. Figure 3 presents the crystal structure of CuPt-ordered InAs\(_{0.50}\)Sb\(_{0.50}\) and the calculated band structure indicating inversion of the \(\Lambda_{4,5}\) and \(\Lambda_{6}\) bands, pointing the semimetallic character.
The quality of InAsSb has improved greatly in the last decade. Significant progress has been made to advance metamorphic buffer layers for LWIR MBE photodetector structures. Different compositionally graded buffer schemes have been studied with the absorber composition \( x \approx 0.60 \), including GaInSb and AlGaInSb graded buffer layers on a GaSb substrate to extend detector’s \( \lambda_{\text{cut-off}} \) [41,42].

In comparison with MBE, MOCVD also allows to grow epitaxial structures on large-area wafers providing higher production throughput (MOCVD is more economical and characterized by less problematical maintenance). However, a disadvantage of this technique is less sharp interfaces, especially in SLs fabrication. More details about InAsSb structures MOCVD growth are included in [20].

3. InAsSb Alloy Properties

Table 2 shows selected properties of semiconductors, along with narrow-gap alloys, used in IR photodetectors manufacturing characterized by zincblende (ZB) or diamond (D) crystal structure. There is a tendency in transition of chemical bond from the covalent group IV-semiconductors to more ionic II–VI materials with the lattice constant increasing when moving across the table from the left to the right. The materials become softer and the chemical bonds become more delicate which results in the quality of the bulk. The semiconductors with a greater share of covalent bonds are more mechanically stable, which contributes to better fabrication. The supreme position of GaAs in optoelectronics and silicon in electronic materials supports this observation. In turn, the energy gap of semiconductors on the right side of the table shows a tendency to have lower values. Higher band-to-band (BtB) absorption provides increased quantum efficiency (QE, \( \eta \)) observed for example in HgCdTe or InSb because of their direct bandgap structure.

A high density of states in the valence band (VB) and conduction band (CB) leading to enhanced absorption of IR radiation and a comparatively low thermal generation rate arise from the direct energy bandgap structure of narrow gap semiconductors applied in the IR detectors. The III–V semiconductors are characterized by much stronger chemical bonds and therefore higher chemical stability compared to HgCdTe, which is important from the producibility viewpoint.

3.1. Energy Gap

The III–V semiconductors are characterized by a ZB structure and direct energy gap at the Brillouin zone centre. The \( \mathbf{k} \cdot \mathbf{p} \) theory explains the shape of the electron and the light mass hole bands. For different semiconductors, the momentum matrix element changes slightly and is typically about \( 9.0 \times 10^{-8} \) eVcm. Therefore, the CB densities of states and the electron effective masses have comparable values for materials with the same energy gap.
Table 2. Fundamental properties of semiconductors used in IR photodetectors manufacturing.

| Semiconductor | Si  | Ge  | GaAs | AlAs | InP   | InGaAs | AlInAs | InAs | GaSb | AlSb | InSb | HgTe | CdTe |
|---------------|-----|-----|------|------|-------|--------|--------|------|------|------|------|------|------|
| Family        | IV  | IV  | III–V| III–V| III–V | III–V  | III–V  | III–V| III–V| III–V| III–V| II–VI| II–VI |
| Lattice constant (Å/structure) | 5.431 | 5.658 | 5.653 | 5.661 | 5.870 | 5.870 | 5.870 | 6.058 | 6.096 | 6.136 | 6.479 | 6.453 | 6.476 |
| Bulk modulus (Gpa) | 98 | 75 | 75 | 74 | 71 | 69 | 66 | 58 | 56 | 55 | 47 | 43 | 42 |
| Band gap (eV) | 1.124 | 0.660 | 1.426 | 2.153 | 1.350 | 0.735 | 0.354 | 0.730 | 1.615 | 0.175 | −0.141 | 1.475 |
| Electron effective mass | 0.26 | 0.39 | 0.067 | 0.29 | 0.077 | 0.041 | 0.024 | 0.042 | 0.14 | 0.014 | 0.028 | 0.090 |
| Hole effective mass | 0.19 | 0.12 | 0.082 (L) | 0.045 (H) | 0.11 (L) | 0.40 (H) | 0.12 (L) | 0.05 (L) | 0.025 (L) | 0.37 (H) | 0.4 | 0.98 | 0.018 (L) | 0.4 |
| Electron mobility (cm²/Vs) | 1450 | 3900 | 8500 | 294 | 5400 | 13800 | 3 × 10⁴ | 5000 | 200 | 8 × 10⁴ | 26500 | 1050 |
| Hole mobility (cm²/Vs) | 505 | 1900 | 400 | 105 | 180 | 500 | 880 | 420 | 800 | 320 | 104 |
| Electron saturation velocity (10⁷ cm/s) | 1.0 | 0.70 | 1.0 | 0.85 | 1.0 | 0.4 | 4.0 | 4.0 |
| Thermal cond. (W/cmK) | 1.31 | 0.31 | 0.5 | 0.7 | 0.27 | 0.4 | 0.7 | 0.15 | 0.06 |
| Relative dielectric constant | 11.9 | 16.0 | 12.8 | 10.0 | 12.5 | 15.1 | 15.7 | 12.0 | 17.9 | 21 | 10.2 |
| Substrate | Si, Ge | GaAs | InP | InAs, GaSb | InSb | CdZnTe, GaAs, Si |
| MW/LW detection mechanism | Heterojunction, internal, photomission | QWIP, QDIP | QWIP | Bulk (MW) | Bulk | Bulk |

D—diamond, ZB—zincblende, id—indirect, d—direct, L—light hole, H—heavy hole, QWIP—quantum well infrared photodetectors, QDIP—quantum dot infrared photodetectors.
The Varshni formula clearly describes a conventional negative temperature coefficient of the energy gap of these materials [43]:

$$E_g(T) = E_0 - \frac{\alpha T^2}{T + \beta'}$$  \hspace{1cm} (1)

where $E_0$ is the bandgap at 0 K, $T$ is the temperature, $\alpha$ and $\beta$ are fitting parameters for a given material. $\beta$ is closely related to the material’s Debye temperature (in Kelvin).

If $T \gg \beta$, then Equation (1) becomes $dE_g(T)/dT \sim -\alpha$. Viña et al. [44] proposed a new expression for temperature dependence of the bandgap energy by taking into account the Bose–Einstein occupation factor [45]

$$E_g(T) = E_0 - \alpha_0 \left( 1 + \frac{2}{\exp(\Theta/T) - 1} \right)$$  \hspace{1cm} (2)

where the parameter $\Theta$ describes the mean frequency of the phonons involved and $\alpha_0$ is the strength of the electron-phonon interaction. While Varshni expression is entirely empirical, the Bose–Einstein retains physical meaning and is more palatable from the theoretical point of view. It is assumed that the Varshni parameter $\beta$ is comparable to the Debye temperature, $\Theta = \langle E_p \rangle/k$, thus the Bose–Einstein average phonon energy can be used to calculate the Debye temperature. If $kT > > \langle E_p \rangle$, the Varshni and Bose–Einstein expressions are equivalent. For lower temperatures in which $kT < \langle E_p \rangle$, the Bose–Einstein formula usually provides a better fit to the experimental results (Ref. [20] discusses in detail the bandgap temperature-dependent parameters).

The dependence $E_g(x,T)$ for InAs$_{1-x}$Sb$_x$ has been experimentally investigated by many research groups since 1964 [22]. The first investigation of the polycrystalline samples optical properties indicated on the nonlinearity of the InAs$_{1-x}$Sb$_x$ bandgap composition dependence, which can be represented by the bowing parameter $C_g$ as

$$E_g(x) = E_{g\text{InSb}}x + E_{g\text{InAs}(1-x)} - C_g(1-x).$$  \hspace{1cm} (3)

Ten different relations $E_g(x,T)$ obtained from measurements over different x-composition values for a wide range of temperatures are presented in [20]. The low-temperature (4–13 K) $\lambda_{\text{cut-off}}$ estimated from $\lambda_{\text{cut-off}} \ [\mu m] = 1.24/E_{g[\text{eV}]}$ changes from 8.4 μm to 12.5 μm. The smallest bandgap of InAs$_{1-x}$Sb$_x$ occurs for $x \approx 0.60$–0.64; however, for such composition lattice-matched binary substrate has not been developed yet. The bandgap expressions from many papers are plotted in Figures 4 and 5 for both high and low temperatures.

![Figure 4. InAs$_{1-x}$Sb$_x$ bandgap energy versus the Sb composition at room temperature. The experimental data is taken with different papers [46–49] as indicated in the legend.](image-url)
At present, $C_g$ value close to 0.7 eV is correct in higher temperature range (Figure 4). The evaluated $E_g(x,T)$ dependence differs from that previously described by Wieder and Clawson in 1973 [46]:

$$E_g(x,T) = 0.411 - \frac{3.4 \times 10^{-4}T^2}{210 + T} - 0.876x + 0.70x^2 + 3.4 \times 10^{-4}xT(1-x).$$

(4)

Initial reports based on experimental data at temperatures above or near 100 K estimate the direct-gap bowing parameter in InAsSb at 0.58–0.6 eV [50]. According to Rogalski and Jóźwikowski’s theoretical considerations, the bowing parameter should tend to move to a higher value of about 0.7 eV [51].

The experimental data from VIGO System collected in recently published paper [47] fits well with this model end equation

$$E_g(x,T) = 0.417 - 1.28 \times 10^{-4}T - 2.6 \times 10^{-7}T^2 - x(C_g + 0.182 + 10^{-9}T^2) + C_gx^2.$$  

(5)

This relation is recommended for InAsSb layers grown on GaAs lattice-mismatched substrates and is valid for both low and high temperatures.

More recent PL studies on unrelaxed MBE-grown InAs$_{1-x}$Sb$_x$ in wide range of composition gave the bowing parameter from 0.83 to 0.938 eV [39,42,49,52]. Until now, the highest bandgap bowing parameter 0.938 eV for low temperature has been reported by Webster et al. [49]. Such large parameter results in the considerable reduction in the smallest bandgap, ~25–75 meV, at low temperature. At room temperature, Webster et al. [49] have obtained much lower bowing parameter of 0.75 eV and developed the temperature-dependent bandgap bowing model. The literature data for InAsSb layers grown by MBE on GaSb fits well with this model end equation

$$E_g(x,T) = 0.417 - 1.28 \times 10^{-4}T - 2.6 \times 10^{-7}T^2 - x(C_g + 0.182 + 10^{-9}T^2) + x^2(C_g - 5.8 \times 10^{-4} + 10^{-7}T^2).$$

(6)

Figure 5 summarizes the experimental data and theoretical predictions of energy gap in temperature range 4–77 K published in different papers. Discrepancies in the $E_g(x,T)$-dependence may result from several factors among others: strains, structural quality of samples, and CuPt-type ordering effect.

Figure 5. InAs$_{1-x}$Sb$_x$ bandgap energy versus the Sb composition at low temperatures. The experimental data are taken with different papers as indicated in the key [39,41,42,49,52–56].

The low energy gap data reported earlier were masked probably by electron filling of the CB, which resulted from background doping due to various degrees of residual strain and relaxation in these
samples. As is marked in Section 2, high quality of unstrained and unrelaxed InAsSb epilayers have been developed by using special graded buffer layers which reduces the substantial difference between the lattice constant of the alloy and the substrate. Electron diffraction patterns of unstrained InAsSb alloys, described in [41,42], showed an ordering-free distribution of group V elements, which indicates that the observed energy gaps of ternary alloys are inherent (both ordering and residual strain effects were eliminated).

The above discussion indicates that the InAsSb bandgap energy is approximately a square function of the composition and shows a weak dependence of the band edge on composition in comparison with HgCdTe (see Figure 6).

![Figure 6. Composition and wavelength diagram of semiconductor material systems.](image)

### 3.2. Bandgap Alignment and Offsets

The SL photodetector’s design process requires the bandgap alignment and offsets for InAsSb in relation to InAs, InSb, and other III–V semiconductor alloys. As is shown in Figure 6, these materials belong to 6.1 Å III–V semiconductor family. They are the most important in proposing new solutions of high-performance IR detectors exhibiting direct energy gaps, high optical absorption, and high design flexibility. This group consists of three alloys having approximately matched lattice constant about 6.1 Å: InAs, GaSb, and AlSb. Their energy gaps are varying in the range starting from 0.417 eV (InAs) to 1.696 eV (AlSb) at lower temperatures [57]. Similarly to other semiconductor compounds, they are chosen as the subject of research due to their heterostructures, in particular combining InAs with the three antimonides (InSb, AlSb, and GaSb) and their alloys. This combination provides a band alignment that is fundamentally different from that of the more thoroughly researched AlGaAs and that band alignment flexibility draws the attention of scientists on the 6.1 Å materials. InAs/GaSb heterojunctions reveal the most unique band alignment, which is defined as the broken gap. The top of the VB of GaSb is located above the bottom of CB of InAs by ~150 meV at the interface. On both sides of the heterointerface, electrons and holes are separated and located in self-consistent QWs, which is related to partial overlapping of the InAs CB with the GaSb VB. Consequently, exceptional tunneling-assisted radiative recombination transitions and novel transport properties are observed. A wide range of different alloys and SLs can be designed due to high versatility of these compounds, with the accessibility of type-I (nested, or straddling), type-II staggered, and type-II broken gap (misaligned) band offsets between the GaSb/AlSb, InAs/AlSb, and InAs/GaSb material pairs, respectively, as illustrated in Figure 7.
Due to different growth conditions causing the variety of residual strain in the materials and due to the discrepancies related to the bandgap measurement methods, the band offsets are historically not easy to estimate. Steenbergen et al. [58] have reviewed the band edge alignment models for InAs-InAs$_{1-x}$Sb$_x$ system and considered different types of heterojunctions. This review is updated in recently published monograph [20]. Figure 8 shows three possible band alignments between InAs and InAs$_{1-x}$Sb$_x$ including two type-II band alignments: with the InAs CB higher in energy than the InAsSb CB and with the InAsSb CB higher in energy than the InAs CB. From these two papers [20,58], the following conclusions could be drawn:

- the type-I alignment is notified for SLs with low Sb compositions and ordering typical for the InAs$_{1-x}$Sbx layer, which is characterized by a lower alloy bandgap than that of a random alloy with equal composition x [39],
- the type-IIa alignment is based on argued conclusions for the reduced mass values of the lowest two transitions given by PL and magneto-transmission measurement results [60,61],
- type-IIb, is chosen for InAs/InAs0.93Sb0.07 SLs on InAs substrates [62].

Investigations carried out during the last several years have revealed the InAsSb band edges versus composition dependence as shown in Figure 9 [20]. The recommended parameters are as follows: InAs/InSb valence band offset (VBO), $-0.59$ eV; the InAsSb bandgap bowing, $0.87$ eV; and the VB bowing, $-0.35$ eV. Moreover, a concept of the band offsets between InAs and InAsSb with slight dependence on temperature was introduced. It follows that the bandgaps are the main temperature-dependent factors.
with the band offsets and VB bowing being distributed by the same fractions in the whole temperature range [49].

![Figure 9. InAs$_{1-x}$Sb$_x$ VB and CB edges and their alignment assuming that $x = 0$ for InAs and $x = 1$ for InSb. As noted by the dashed arrow, longer wavelengths than InAsSb can be reached with T2SLs grown on a virtual substrate In$_{0.54}$Sb$_{0.46}$ (after [20,63]). Bandgap bowing, $C_g$, has been distributed to the VB, $C_v$, and CB as shown according to the references [49,64]. The dash-dot curves represent the suggested bandgap and VB bowing equal 0.87 and $-0.35$ eV, respectively. The solid arrow points the composition of In$_{0.91}$Sb$_{0.09}$ lattice-matched to GaSb substrates.](image)

The InAsSb properties are direct consequence of the energy band structure. Table 3 contains some material parameters of the InAs, InSb, GaSb, and InAs$_{0.35}$Sb$_{0.65}$ semiconductors [3,65]. Between them, InSb has been investigated most broadly.

3.3. Transport Properties

Virtual crystal approximation (VCA) describes the electronic properties of ternary alloys according to the approach [66], where the disordered alloy is modeled by an ideal crystal with an average potential at each sublattice site. Linear interpolation of the potentials of corresponding binary semiconductors is realized by the potential averaging. The nonlinearity of composition dependence versus the energy gap (see Equation (3)) is calculated based on VCA model because of the nonlinear relationship of the band energies against the crystal potential. In order to explain the large bandgap bowing, the VCA has to be modified by adding the random potential due to fluctuations of the alloy composition [67,68]. Presently, it is commonly assumed that the material disorder influences not only the large bandgap bowing in the InAsSb alloy but also shapes the composition dependence on fundamental parameters such as mobilities or carrier effective masses.

The conduction-valence-band mixing theory [23] has been implemented to obtain a good agreement between experimental room temperature effective masses and calculations, as reported by Rogalski and Jóźwikowski [51] (see Figure 10). More recently published low-temperature data, especially for mid-composition range, indicates lower electron effective masses. Estimated negative bowing parameter for the electron effective mass is $C_m = 0.038$ and is slightly less than expected from the Kane model ($C_m = 0.045$), reaching the lowest effective mass (0.0082$m_0$ at $x = 0.63$ and 4 K) ever reported for III–V semiconductors. A possible reason for the effective mass value discrepancy, shown in Figure 10, is the alloy disorder causing the random potential distribution which disturbs the CB and VB states.
The electrical properties of semiconductor compounds and their dependence on composition are very important for device design with the optimal performance. However, it is difficult to control carrier concentration because insulating or semi-insulating substrates for InAsSb epitaxial layers are generally not available. Parallel conduction layers such as the substrate and/or the interface with the

### Table 3. Physical properties of narrow gap III–V alloys.

| Lattice structure | Ti(K) | InAs | InSb | GaSb | InAs_{0.35}Sb_{0.65} |
|-------------------|-------|------|------|------|----------------------|
| Lattice constant a (nm) | 300 300 80 300 1210 | 0.60584 | 0.647877 | 0.6094 | 0.636 |
| Thermal expansion coefficient α (10^{-6} K^{-1}) | 300 | 80 | 5.02 | 5.04 | 6.50 | 6.02 |
| Density ρ (g/cm^3) | 300 | 5.68 | 5.7751 | 5.61 |
| Melting point T_m (K) | 1210 | 803 | 985 |
| Energy gap E_g (eV) | 4.2 80 300 | 0.42 0.414 0.359 | 0.2357 0.228 0.180 | 0.822 0.226 0.725 | 0.138 0.136 0.100 |
| Thermal coefficient of E_g | 100–300 | 0.65 | 0.43 | 0.28 | 0.41 | 0.10 |
| Effective masses: | | | | | |
| m_e/m | 4.2 300 4.2 | 0.023 0.022 0.026 | 0.0145 0.0116 0.0149 | 0.042 0.0101 |
| m_h/m | 4.2 300 | 0.43 | 0.41 | 0.28 | 0.41 |
| m_{le}/m | | 4.2 | 0.43 | 0.28 | 0.41 |
| Momentum matrix element P (eV cm) | | | | | |
| | | 9.2 \times 10^{-8} | 9.4 \times 10^{-8} |
| Mobilities: | 77 | 300 77 300 | 8 \times 10^4 3 \times 10^4 8 \times 10^4 3 \times 10^4 1 \times 10^4 | 10^6 | 5 \times 10^3 | 5 \times 10^4 |
| \mu_e (cm^2/Vs) | | | | | |
| \mu_h (cm^2/Vs) | | | | | |
| Intrinsic carrier concentration n_i (cm^{-3}) | 77 200 300 | 6.5 \times 10^3 7.8 \times 10^{12} 9.3 \times 10^{14} | 2.6 \times 10^9 9.1 \times 10^{14} 1.9 \times 10^{16} | 2.0 \times 10^{13} 8.6 \times 10^{14} 4.1 \times 10^{16} |
| Refractive index n_i | | 3.44 | 3.96 | 3.8 |
| Static dielectric constant ε_s | | 14.5 | 17.9 | 15.7 |
| High frequency dielectric constant ε_{oo} | | 11.6 | 16.8 | 14.4 |
| Optical phonons: | | | | | |
| LO (cm^{-1}) | | 242 | 193 | 220 |
| TO (cm^{-1}) | | 220 | 185 | 200 |

![Figure 10. Effective mass of electron versus composition for InAs_{1-x}Sbx. The experimental data is taken from [51,56].](image)

The electrical properties of semiconductor compounds and their dependence on composition are very important for device design with the optimal performance. However, it is difficult to control carrier concentration because insulating or semi-insulating substrates for InAsSb epitaxial layers are generally not available. Parallel conduction layers such as the substrate and/or the interface with the
substrate presence make Hall effect measurements difficult to interpret. Moreover, it has been well known from the beginning of the 1970s [69,70] that narrow gap III–V semiconductors, especially InAs, creates surface electron accumulation layers. For example, Figure 11 presents band edges of p-type InAs [69] and InAsSb [71] with surface electron accumulation. There is strong electron accumulation on the InAsSb side of InAsSb/GaSb heterojunction due to the staggered type II band alignment. As is shown in Figure 11b, the surface states cause the Fermi level to be pinned above the CB creating a parallel conduction path to the epilayer. In this case, the accurate determination of the carrier concentration and mobility with standard Hall measurements is challenging.

![Figure 11](image)

**Figure 11.** Band edges of p-type InAs and InAsSb with surface electron accumulation: (a) InAs with surface inversion layer; the electrostatic potential has the value $\Phi_s$ at surface and $\Phi_b$ in the bulk, (b) InAsSb/AlAsSb heterojunction with depicting surface electrons, epilayer, and interface with AlAsSb.

The above-mentioned effect should be taken into account to analyze electrical carrier concentration and mobility measurements. A popular two-layer model has been often used to separate the surface accumulation and epilayer properties. Another approach is applied in the case of more complicated structures like that shown in Figure 11b. Mobility spectrum analysis and/or variable temperature and magnetic field-dependent transport analysis with multicarrier fitting models have been applied to segregate the epilayer and surface and interface layer characteristics [71].

The variation of InAs$_{1-x}$Sb$_x$ electrical properties depend on the composition. Similarly to binary compounds, ternary alloys also have the surface accumulation layers and interface layers with the buffers or substrates. Special attention is required in estimates of carrier concentrations in selected device regions. More information on that topic can be found in [20].

As a popular donor dopant in InAs, silicon is used. Beryllium and zinc have been used for InAs and InSb photodiode fabrications for conversion of n-type materials to p-type ones [5]. Silicon behavior in InSb is amphoteric; in pulled crystals results in p-type conductivity for temperatures $>400$ °C, however for MBE growth epilayers results in n-type when the growth temperature is $>350$ °C, but for growth temperatures $<320$ °C gives donor conductivity [20]. Germanium is used as an acceptor dopant in InSb but behaves as a donor in InAs for LPE-grown epilayers [72].

Rogalski and Jóźwikowski have calculated the InAs$_{1-x}$Sb$_x$ intrinsic carrier concentration for the conditions $0 \leq x \leq 1$ and $50 \leq T \leq 300$ K. In order to obtain a good agreement between experimental values and calculations, the conduction-valence-band mixing was taken into account [51]:

\[
n_i = \left(1.35 + 8.50x + 4.22 \times 10^{-3}T - 1.53 \times 10^{-3}xT - 6.73x^2\right) \times 10^{14}T^{3/2}E_s^{3/4} \exp\left(-\frac{E_g}{2kT}\right),
\]

where $n_i$ is in cm$^{-3}$ and $k$ is the Boltzmann constant. The maximum of $n_i$ occurs at $x \approx 0.63$ which corresponds to the minimum energy gap for a given temperature.

The CB density of states is low due to the very small effective mass of electrons and filling the available band states can be realized by doping shifting the absorption edge to shorter wavelengths.
This phenomenon has been recognized as the Burstein-Moss (BM) effect, which is shown in Figure 12 for n-type InAs$_{0.855}$Sb$_{0.145}$ epilayer with residual background donor concentration of $3 \times 10^{16}$ cm$^{-3}$.

The theoretically predicted absorption was calculated using the Kane model, including the BM shift according to Anderson theory [73]. As is shown, the experimental spectrum is in proper agreement with the theoretical absorption simulation at 20 K and 300 K. The bandgap energy values measured with IR transmission spectroscopy agree with those obtained from PL peaks, particularly at low temperatures.

The first measurements of transport properties of n-type InAsSb alloys were performed on samples prepared in the late 1960s by various freezing and annealing techniques [74,75]. The properties of high-quality InAsSb epitaxial layers with $x < 0.35$ fabricated by LPE are comparable to those of pure InAs (when $n = 2 \times 10^{16}$ cm$^{-3}$, typical mobilities are 30,000 cm$^2$/Vs at 300 K and 50,000 cm$^2$/Vs at 77 K). For InSb-rich alloys with $x \geq 0.90$, typical mobilities are 60,000 cm$^2$/Vs at 300 K. When As is added to InAs$_{1-x}$Sb$_x$ alloys, the residual background concentration has increased to a low $10^{16}$ cm$^{-3}$; in turn the mobility first increased and then dropped by a factor of 1.5 to 2 as the temperature decreases from 300 to 77 K. At the present stage of MBE-growth development, the background electron concentration in InAs$_{1-x}$Sb$_x$ with 40% Sb at 77 K is as low as 1.5 $\times$ $10^{15}$ cm$^{-3}$ [39].

The carrier mobility in the III–V binary compounds is primarily limited by scattering due to optical phonons, and ionized impurities with a contribution from alloy disorder in the corresponding ternary alloy. The last kind of scattering mechanism is so-called alloy scattering. Their relative contributions depend on temperature, carrier concentration, compensation, and alloy composition. Acoustic phonon and deformation potential, optical deformation potential, and neutral impurity scattering are minor effects under most conditions. The total carrier mobility $\mu_{tot}$ in alloy InAs$_{1-x}$Sb$_x$ can be simply expressed as [27]:

$$\frac{1}{\mu_{tot}(x)} = \frac{1 - x}{\mu_{totInSb}} + \frac{x}{\mu_{totInAs}} + \frac{x(1 - x)}{C_\mu},$$

(8)

The first two terms of Equation (8) result from the linear interpolation scheme and the third term reflects the effects of alloying.

The InAs$_{1-x}$Sb$_x$ mobility composition dependence is illustrated in Figure 13 at 77 and 300 K for carrier concentrations of experimental significance [76,77]. The component mobilities are shown for carrier concentrations of $n = 10^{15}$, $10^{16}$ and $10^{17}$ cm$^{-3}$ at 77 K, and $10^{16}$, $10^{17}$ cm$^{-3}$ at 300 K. A decrease in impurity concentration causes continuous increase of carrier mobility throughout the temperature range. The significance of alloy scattering is visible when temperature and carrier concentration decrease as the respective contributions from polar optical phonons and ionized impurities are reduced, until it remains as the dominant scatterer across most of the compositional range for low carrier concentrations (below $10^{15}$ cm$^{-3}$) and low temperatures.
Figure 13. InAs$_{1-x}$Sb$_x$ mobility composition dependence at 77 K. The solid lines are the calculated mobilities for selected carrier concentrations, and the dashed lines are the component mobilities determined for a carrier concentration of 10$^{15}$ cm$^{-3}$ [77]. Experimental data is marked for epitaxial layers with carrier concentrations between 10$^{15}$ to 10$^{16}$ cm$^{-3}$. Scattering legend: $\mu_{\text{op}}$-optical phonons, $\mu_{\text{i}}$-ionized impurity, $\mu_{\text{alloy}}$-alloy disorder.

Egan et al. have calculated the electron mobility of InAsSb by considering all the possible scattering mechanisms: impurities, acoustic phonons, optical phonons, alloy scattering, and dislocations [77]. Comparison with experiment confirms that dislocation scattering has a strong effect on transport, while alloy scattering limits mobility in ternary samples grown with a minimum of defects (see Figure 14).

Figure 14. InAs$_{1-x}$Sb$_x$ electron mobility composition dependence for room-temperature (solid lines [77]) and carrier concentration of 10$^{17}$ cm$^{-3}$ with a compensation ratio of 0.2 and the effects of a dislocation density of 3.8 $\times$ 10$^8$ cm$^{-3}$. The experimental data is taken after [20,23] for samples with electron concentrations above 10$^{16}$ cm$^{-3}$.

The next figure (Figure 15) presents the modeled InAs$_{0.80}$Sb$_{0.20}$ electron and hole mobilities temperature dependence with low residual background doping [78]. As is shown, the mobility ratio $b = \mu_e/\mu_h$ assumes $\sim$10$^2$, similarly to InSb.
where the electron density in the active layer is reduced below its equilibrium level, the internal annihilation of electron-hole pairs with emission of photons makes an inverse process. Three predominant mechanisms in this type of band structure are presented. They are determined by the steady-state concentration of carriers in semiconductors based on the optical (kinetics of photogenerated carriers) and thermal excitation. A wide study of G-R processes in semiconductors is reported in the literature (see for example [79–81]). We describe here only selected carrier lifetime results directly related to the photodetector performance. Assuming bulk processes only, there are three main thermal G-R processes to be considered in the narrow bandgap semiconductors: SRH, radiative, and Auger.

The statistical theory for the G-R processes via intermediate centers was developed first by Shockley and Read [82], and Hall [83]. It may be reduced by decreasing foreign impurities and native defects concentrations, which can be achieved by low-temperature growth and by progress in the material’s development. Since the SRH does not represent a fundamental limit to photodetector performance (it can be reduced with progress toward purer and higher-quality material), a significant research effort is still needed.

Absorption of internally generated photons causes the radiative generation of charge carriers. The annihilation of electron-hole pairs with emission of photons makes an inverse process. Internal radiative processes are the fundamental factor limiting the detector performance. However, critical re-examination of the radiative mechanism contribution in the IR detection has been presented in the literature [84,85]. Humphreys [84] indicated that due to photon reabsorption (PR), the radiative lifetime is highly extended and depends on the semiconductor geometry. Under reverse bias operation where the electron density in the active layer is reduced below its equilibrium level, the internal radiative generation could be suppressed [86]. In spite of the fundamental nature, the radiative processes do not limit significantly the ultimate performance of IR detectors, especially in the LWIR range [87].

In high-quality narrow gap semiconductors such as InSb and Hg$_{1-x}$Cd$_x$Te, Auger mechanisms determine G-R processes at near room temperatures. The Auger mechanisms in InSb-like band structure materials are classified into 10 photonless mechanisms according to the related bands. Figure 16 presents the three predominant mechanisms in the case of this type of band structure. They are characterized by the largest combined density of states and smallest threshold ($E_F \approx E_g$).
The conduction band/heavy-hole band/conduction band (CHCC) recombination mechanism (also labeled Auger 1) prevails in n-type material and involves two electrons and a heavy hole. For p-type material the conduction band/heavy-hole band/light-hole band (CHLH) process (marked as Auger 7) is characteristic under condition that the spin split-off band can be neglected. Auger transition probability through the conduction band/heavy-hole band/spin split-off band mechanism (labeled CHSH or Auger S hereafter) can be ignored for semiconductors such as InSb and HgCdTe where the spin split-off energy (Δ) is much larger than the bandgap energy E_g and the Auger 7 transition is typical.

The spin split-off band is even more essential than the light-hole band for the direct-bandgap materials, particularly when the bandgap energy E_g approaches the spin-orbit splitting Δ (as in the case of InAs and InAsSb). The carrier lifetimes in binary compounds (InSb and InAs) are discussed in detail in chapter 2 monograph Antimonide-Based Infrared Detectors. A New Perspective [88].

Figure 17 summarizes the InAs_{1-x}Sbx spin-orbit-splitting energy, Δ, composition dependence at 10 K. Experimental results of Δ(x) dependence published in 1972 by Van Vechten et al. [75] differ more strongly than that presented recently in the paper by Cripps et al. [89]. The Δ parameter versus x (Sb fraction) characteristic does not show bowing clearly. A good approximation can be calculated with

\[ \Delta(x) = 0.81x + 0.371(1-x) + 0.165x(1-x) \]  

The above formula allows for the calculation of the spin-orbit-splitting bandgap energy which stays in a good agreement with experimental results and does not depend on temperature.

**Figure 16.** The three BtB Auger recombination processes. Arrows indicate electron transitions; ●, occupied state; ○, unoccupied state.
For slight deviation from equilibrium and for nondegenerate materials with equilibrium carrier concentration $n_0$ (or $p_0$), the carrier lifetimes can be calculated using the intrinsic carrier lifetime and $n_0/n_i$ (or $p_0/n_i$) ratio [65,79]:

for radiative recombination:

$$\tau_{R}^{j} = \frac{2\tau_{R}^{i}}{(n_0/n_i) + (n_i/n_0)} = \frac{2\tau_{R}^{i}}{(p_0/n_i) + (n_i/p_0)}.$$  \hspace{1cm} (10)

for Auger recombination:

$$\tau_{A1} = \frac{2\tau_{A1}^{i}}{1 + (n_0/n_i)^2}, \quad \tau_{A7} = \frac{2\tau_{A7}^{i}}{1 + (p_0/n_i)^2}, \quad \tau_{AS} = \frac{2\tau_{AS}^{i}}{1 + (p_0/n_i)^2}.$$  \hspace{1cm} (11)

The InAs$_{1-x}$Sb$_x$ Auger and radiative recombination carrier lifetime for the composition range $0 \leq x \leq 1$ and the temperature range 77–300 K was estimated by Rogalski and Orman [90]. Radiative recombination determines the carrier lifetime in the low temperature range. At higher temperature, the Auger 1 process is dominant in n-type InAs$_{1-x}$Sb$_x$ but in p-type material, for the composition range $0 \leq x \leq 0.3$, Auger 7 and Auger S processes compete while for $x > 0.3$, the Auger 7 process dominates. Different recombination mechanisms in InAs$_{0.35}$Sb$_{0.65}$, operating at the longest $\lambda_{cut-off}$ (with $E_G$ close to 0.1 eV at room temperature) are also discussed in [90].

The dependence of $\tau_{R}^{i}, \tau_{A1}^{i}, \tau_{A7}^{i}$ and $\tau_{AS}^{i}$ on the composition $x$ in InAs$_{1-x}$Sb$_x$ at room temperature is presented in Figure 18. The effect of the Auger S process was considered as accurate for $0 \leq x \leq 0.3$ according to the out-of-date carrier lifetime calculation [90]. The predicted contribution of Auger S process is dominant in InAs$_{1-x}$Sb$_x$ semiconductors close to InAs (for $0 \leq x \leq 0.15$) taking more recently estimated dependence of $\Delta(x)$ [89]. It follows that InAsSb photodiodes have lower Auger S nonradiative losses than were reported previously [91]. This adjustment has substantial implications on the p-type InAsSb-based devices’ design.

![Figure 18. Intrinsic InAs$_{1-x}$Sb$_x$ Auger and radiative recombination carrier lifetimes at room temperature. $\tau_{S1}^{i}$ is calculated assuming spin-orbit-splitting bandgap energies after [75,89] (after [91]).](image)

Generally, in III–V semiconductors the carrier lifetime is not limited by the BtB recombination but by SRH mechanism. Two regions are of interest:

- the neutral or diffusion region of the detector, and
- the depletion region of the detector.
As discussed in many papers, there are two types of possible SRH defects—neutral centers at the intrinsic level with carrier lifetimes $\tau_{ne} \sim \tau_{i0}$ and charged centers at some specific energy level in which there is a large asymmetry in the lifetimes associated with electrons and holes.

In comparison with II–VI HgCdTe, the reported SRH lifetimes in the majority of III–V binary alloys are not impressive. For example, the best values for InSb indicate that $\tau_{SRH}$ is about 400 ns for LPE-grown material [92]. It is interesting to stress that this value has not been improved over the last 60 years, and no response defect has been identified. Similar lifetime was reported for MBE grown InAs/InAs$_{0.72}$Sb$_{0.28}$ T2SLs. Assuming this carrier lifetime value, Kinch modeled the minority carrier lifetime and diffusion length versus temperature for holes in n-type InAs$_{0.81}$Sb$_{0.19}$ with doping levels $10^{15}$ and $10^{16}$ cm$^{-3}$—see Figure 19 [78]. These values are approximately three orders of magnitude smaller than those in HgCdTe with similar bandgaps.

![Figure 19. Simulated minority carrier lifetime and diffusion length versus temperature for holes in n-type InAs$_{0.81}$Sb$_{0.19}$ with doping levels $10^{15}$ and $10^{16}$ cm$^{-3}$ and $\tau_{SRH} = 400$ ns.](image)

### 3.5. Other Properties

The optical properties of InAsSb are determined by a nonparabolic three-band model. Dobbelare et al. [93] have studied IR absorption spectra of Si-doped MBE epitaxial layers grown on Si and GaAs substrates. The absorption coefficient was measured in the wavelength range 3–12 μm and the experimental data were fit by an analytical formula that was derived from the Kane band model. Excellent agreement between the calculations and the measurements was observed (Figure 20). Due to the fact that the measurements were carried out for InAs$_{1-x}$Sb$_x$ with $x \approx 0.70$, a large shift of the absorption edge with Si doping was observed due to low density of states in the CB. This has been referred to the BM effect.

Comparable values of absorption coefficients for bulk HgCdTe [94,95] and InAsSb [49,96] are reported (see Figure 21), proving relatively minor differences between densities of states in bulk semiconductors with the same energy gap and the optical matrix elements.

The Raman spectroscopy is a well-known technique to understand the lattice vibrations of various material systems. The first Raman spectroscopy study of InAs$_{1-x}$Sb$_x$ on InAs and InSb substrates was performed by Cherng et al. [97]. They reported “one-mode” phonon behavior for $x \leq 0.6$ and “two-mode” phonon behavior for larger values. Then, a similar study for InAsSb grown with different composition on GaAs substrate by MBE was carried out in [98,99], and “two-mode” phonon behavior was observed. Along with the Sb composition increasing in InAs-like longitudinal-optical (LO), transverse-optical (TO) phonon peaks reveal a blue and red-shift occurring for InSb-like LO phonon peak—see Figure 22. Recently, Grodecki et al. [100] suggested that spectra in the region 20–160 cm$^{-1}$ are
caused by zone folding in phonon dispersion curves as a result of CuPt unit cell of InAsSb being twice as large as typical ZB unit cell.

Figure 20. Absorption spectra for InAs$_{0.30}$Sb$_{0.70}$ grown on GaAs substrates at 300 and 77 K with electron concentrations (a) $10^{17}$ cm$^{-3}$ and (b) $2 \times 10^{18}$ cm$^{-3}$. Solid lines present theoretical fitting (after [93]).

Figure 21. Intrinsic absorption coefficient for Hg$_{1-x}$Cd$_x$Te, InSb, and lattice-matched InAs$_{0.911}$Sb$_{0.09}$ and InAs/InAsSb SLs at room temperature.

Figure 22. InAs$_{1-x}$Sb$_x$ Raman frequency dependence versus composition (after [99]).
4. Properties of InAs/InAsSb Superlattices

Ting et al. published simple theoretical considerations on the fundamental T2SLs parameters [101]. Their properties differ strongly from those of constituent layers.

InAs/InAsSb SLs have been less researched than InAs/GaSb SL and they are in the early stage of development. Because of only two common elements (In and As) in SL layers and rather uncomplicated interface structure with Sb-shifting elements, the InAs/InAsSb SLs growth follows with a better controllability and easier manufacturability. It appears that InAs/InAsSb SLs are more flexible in the optimization of device performance. In order to reduce the tension between the SL layers and the substrate on which the SL is deposited, a suitable selection of the buffer layer is required in which the tension between the SL and the buffer layer is averaged. Usually, in order to fabricate the defect-free buffer layer, the interfacial misfit array (IMF) technique is applied [102]. By producing the strain-balanced SL to the buffer layer, the influence of the strains and dislocations on the device performance is eliminated.

The carrier localization effect is noticed in T2SLs at low temperature because of local impurities and spatial variations in the CB and VB potentials, layer thickness variation, or composition variation. These local potentials result in low-energy tail states (lower than the bandgap), which can trap carriers. The presence of carrier localization is supported by observation of extremely long carrier lifetimes and PL peak blue shifts at low temperatures [20,103]. The InAs/InAsSb interface disorder was found to be the reason for the carrier localization [104]. Therefore, the greater effect from interface disorder and deeper localization potentials are observed in smaller SL periods. The carrier localization effects are not observed at higher temperatures (>100 K).

4.1. Bandgap Energy and Effective Masses

Since InAs is not lattice-matched to InAsSb, the changing of the layer thickness requires a good understanding of the strain contribution on the material quality. Figure 23 shows the SLs InAs/InAs0.62Sb0.38 band structure for unstrained and strained conditions. Under the influence of strain, an electron affinity changes and the VB splits on the heavy and light holes’ sub-bands.

![Figure 23. The energy band profiles of unstrained and strained InAs/InAs0.62Sb0.38 SLs (after [105]).](image)

The basic difference in profiles of the CB and VB in the InAs/GaSb and InAs/InAsSb T2SLs is presented in Figure 24. The band alignment of T2SL is causing a state in which the energy bandgap of the SLs can be adjusted to the configuration, either a semimetal or a narrow bandgap semiconductor material.
Transitions between electron and hole bands are spatially indirect. The first electron miniband (C₁) is more sensitive to layer thickness than first heavy hole state (HH₁) because of the large value of the heavy-hole mass.

Below are emphasized distinctions in the fundamental properties of InAs/GaSb and InAs/InAsSb SLs:

- both type of T2SLs are based on nearly-lattice-matched III-V semiconductors and provide a large range of tunability in λ_{cut-off},
- in T2SLs the electron and hole wave functions are located in separate layers,
- the resulting energy gap is determined by the transition energy between the HH₁ and the C₁ and depends upon the layer thicknesses and interface compositions,
- the band offsets in conduction (ΔE_c) and valence (ΔE_v) bands in the InAs/InAsSb SL (ΔE_c ~142 meV, ΔE_v ~226 meV) are much smaller as compared to InAs/GaSb SL (ΔE_c ~930 meV, ΔE_v ~510 meV),
- a much larger broken gap of the InAs/GaSb SLs makes it easier to reach small SL bandgaps,
- as the period increases, the λ_{cut-off} of the InAs/GaSb SL increases much faster than in the case of the InAs/InAsSb SL. Shorter period of InAs/GaSb SL gives the same λ_{cut-off} as for InAs/InAsSb SL.

Figure 25 shows the theoretically estimated the λ_{cut-off} versus SL period for strain balanced (with respect to the GaSb substrate) InAs/GaSb and InAs/InAsSb T2SLs. In order to achieve strain balance between InAs/GaSb SLs layers, InAsSb interfacial MLs are inserted between the InAs and GaSb layers. As is shown in Figure 25, the (m,n)A/B denotation means SLs with MLs of material “A” and MLs of material “B” in each period of (m + n) MLs.

![Figure 24. Bandgap diagram for (a) InAs/GaSb and (b) InAs/InAsSb T2SLs (after [106]).](image)

![Figure 25. Theoretically estimated λ_{cut-off} for (2n,n)InAs/GaSb, (m,7)InAs/GaSb, (3n,n)InAs/InAs_{0.6}Sb_{0.4}, and (4n,n)InAs/InAs_{0.5}Sb_{0.5} SLs at 100 K (all on GaSb substrate) versus SLs period in MLs (after [107]).](image)
Due to the considerably larger effective mass along the growth direction than those in bulk semiconductors, particularly in LWIR range, T2SLs are characterized by a low mobility, a short diffusion length, and low collection efficiency in n-type devices. Figure 26 shows a comparison of the calculated electron and hole effective masses for bulk InAsSb and both InAs/GaSb and InAs/InAsSb T2SLs versus $\lambda_{\text{cut-off}}$ [107,108]. Generally, the rate of increase for both $m_{e,z}^*$ and $m_{h,z}^*$ versus $\lambda_{\text{cut-off}}$ is faster in SL requiring longer periods to reach the same $\lambda_{\text{cut-off}}$. An increase of the SL period to reach longer $\lambda_{\text{cut-off}}$ leads to the weaker coupling between wave functions and in consequence results in larger effective masses. The relative insensitivity of the InAs/GaSb T2SL effective mass along the growth direction, $m_{e,z}^*$, versus $\lambda_{\text{cut-off}}$ results from the fact that the HH1 state is located within the broken gap energy range. We can notice that in properly designed T2SLs, the electron effective masses are comparable to those in bulk material. However, in the LWIR region the hole effective masses along the growth direction of T2SLs are noticeably larger than those in bulk. In comparison with HgCdTe with the bandgap energy $E_g \approx 0.1 \text{ eV}$ and $m^*_e = 0.009m_0$, the electron effective mass of T2SLs is larger. Comparison of both types of the SLs in MWIR and LWIR ranges indicates that the InAs/GaSb T2SL would have smaller $m_{h,z}^*$ in the very-long wavelength (VLWIR).

Figure 26. Electron (a) and hole (b) effective masses along the growth direction versus $\lambda_{\text{cut-off}}$ for different material systems: free-standing bulk InAsSb with varying Sb composition $x$, (m,7)-InAs/GaSb T2SL, pseudomorphic (m,n)-InAs/InAs$_{0.6}$Sb$_{0.4}$ T2SL on GaSb substrate, pseudomorphic (m,n)-InAs/InAs$_{0.5}$Sb$_{0.5}$ T2SL on GaSb substrate and (m,n)-InAs/InAs$_{0.5}$Sb$_{0.5}$ T2SL on a metamorphic substrate with lattice constant 0.4% larger than that for GaSb (after [107]).

4.2. Absorption Coefficient

The InAs/InAsSb T2SL with an average lattice constant matched to GaSb have significantly lower absorption than InAs/GaSb one (see Figures 21 and 27). It is related to the overlapping of the electron-hole wave functions, which in the case of the InAs/InAsSb T2SL occurs primarily in the well for holes having a relatively small thickness throughout the SL period. The InAs/GaSb SLs have the more favorable oscillator strengths and absorption coefficients than InAs/InAsSb, especially in VLWIR spectral range. However, due to weaker confinement of the CB, the more delocalized electron wave functions help somewhat in enhancement of the absorption properties of the InAs/InAsSb T2SLs. Generally, a shorter SL period provides a larger absorption coefficient due to a better wave function overlap and greater oscillator strength.

In general, the absorption coefficient near $\lambda_{\text{cut-off}}$ is weaker for the InAs/InAsSb SL than for the InAs/GaSb SL. From the other side (see Figure 21), the absorption coefficient of HgCdTe is stronger than those for both T2SLs. In addition, theoretical estimates of the absorption spectra presented by Vurgaftman et al. [109] for LWIR T2SLs and bulk materials (HgCdTe and InAsSb) support above observations-see Figure 27b. In addition, these estimates show that only for the
small-period metamorphic In$_{1-x}$Sb$_x$/In$_{1-y}$As$_y$ SLs, its value of absorption coefficient is comparable with bulk materials.

Figure 27a demonstrates experimental and theoretical absorption coefficient for InAs/GaSb and InAs/InAsSb T2SLs [110,111]. Theoretical calculations in Figure 27a reproduce well the experimental absorption coefficient for 12.8ML/12.8ML InAs/InAsSb SL and for the 8.4ML/13.7ML InAs/GaSb SL. Visible for the 8.4ML/13.7ML InAs/GaSb SL strong peak at a wavelength of about 2.6 $\mu$m is the result of the zone boundary HH$_2 \rightarrow$ C$_1$ transitions. Another slightly weaker peak at wavelength of about 3.4 $\mu$m is due to the zone centre LH$_1 \rightarrow$ C$_1$ transition.

![Figure 27](image-url)

**Figure 27.** Experimental and theoretical absorption spectra for MWIR InAs/GaSb and InAs/InAsSb T2SLs at 77 K (a) (adapted after [110]) and LWIR HgCdTe and different III-V material systems at 80 K (b). (b) presents only calculated interband absorption coefficients for bulk InAs$_{0.60}$Sb$_{0.40}$, Hg$_{0.76}$Cd$_{0.24}$Te, and T2SLs: 42Å InAs/21Å GaSb, 96Å InAs/29Å InAs$_{0.61}$Sb$_{0.39}$ and 11Å InAs$_{0.66}$Sb$_{0.34}$/12Å InAs$_{0.36}$/Sb$_{0.64}$ metamorphic.

### 4.3. Carrier Lifetimes

Apart from simpler manufacturability of InAs/InAsSb T2SLs in comparison with InAs/GaSb system, the interest in InAs/InAsSb SLs stemmed mainly due to the lack of restriction on the carrier lifetime resulting from the Ga presence in the InAs/GaSb SL.

In general, the carrier lifetime is enhanced in T2SLs due to suppression of Auger mechanisms resulting from the separation of electrons and holes. Since the optical transitions occur spatially, the optical matrix element for BTB transitions is relatively small. Theoretical calculations show that Auger recombination rates in T2SLs are suppressed by several orders in comparison to the bulk HgCdTe optimized at the same $\lambda_{\text{cut-off}}$. Nevertheless, these theoretical considerations have not been experimentally confirmed yet. In practical devices based on III-V material systems, more active SRH centers are observed, compared to those of HgCdTe ternary alloys, resulting in lower carrier lifetime. What is worth noting is that the SRH carrier lifetime in InSb has been on the same level since the beginning of research in the 1950s. The situation is more complex in the case of T2SLs.

In accordance with the statistical theory of the SRH mechanism, the shortest SRH lifetime occurs through the trap centers located approximately in the intrinsic energy level in the bandgap. The location of the Fermi level has an impact on the location of the energy level associated with native defects. For example, the Fermi level in bulk GaSb is located either in the mid-gap or near the VB edge, while in bulk InAs is located above the CB edge. Thus, the trap level located in the mid-gap of GaSb promotes the SRH recombination, whereas in InAs trap centers are inactive for the SRH process. For this reason, the carrier’s lifetime in bulk InAs is longer than in bulk GaSb. It can be assumed that the native defects associated with the GaSb layer (SRH centers has been attributed to the presence of gallium) in InAs/GaSb T2SL are responsible for the SRH-limited minority carrier lifetime. The gallium-free (Ga-free)
SLs show much longer carrier lifetimes, up to 10 µs for undoped material in MWIR region [112,113]. The minority carrier lifetime in InAs/InAsSb SLs increases versus Sb composition and lower thickness of InAsSb layer.

At low temperature, the carrier lifetime in the InAs/InAsSb SLs is enhanced by the carrier localization effect. The localized trap states tend to be in resonance with the CB rather than in the bandgap. As shown in the inset of Figure 28, the minority carrier lifetime for MWIR T2SLs decreases to 1.26 µs versus temperature and for T = 77 K is an order of magnitude shorter than that at 15 K. It is suggested that at 15 K the photo-generated electron-hole pairs recombine through spatially separated localized states. A long minority carrier lifetime of 12.8 µs is due to a small overlapping of the wave functions.

![Figure 28](image)

**Figure 28.** Time-resolved PL decays of samples A (bulk InAsSb) and B (MWIR InAs/InAsSb T2SL) at 15 K. The minority carrier lifetimes of 1.24 µs and 12.8 µs are extracted from the single exponential decays, respectively. The inset shows the carrier lifetime of sample B versus temperature (after [104]).

In devices operating at higher temperatures (typically above 80 K), no deleterious effects from carrier localization are observed. However, for detectors with low background doping, operating at low temperatures (typically below 80 K) and under low incident light intensities, carrier localization may be visible; e.g., reduced carrier collection efficiency resulting in lower QE.

Figure 29 presents the measured and modeled hole minority carrier lifetime versus inverse temperature for n-type InAs/InAs$_{0.54}$Sb$_{0.46}$ T2SL with doping concentration $N_d = 5 \times 10^{14}$ cm$^{-3}$ [114]. In the extrinsic region, $(N_d > n_i)$, the lifetime is dominated by the SRH lifetime, $\tau_{SRH}$. Taking the measured lifetime to be 4.5 µs and the radiative lifetime of 9.3 µs, the calculated SRH lifetime is 8.7 µs, if recombination centre is located at the intrinsic Fermi level. The Auger hole lifetime curve is presented for overlap integral $|F_1F_2| = 0.22$.

![Figure 29](image)

**Figure 29.** Hole minority carrier lifetime versus inverse temperature for a n-type MWIR InAs/InAs$_{0.54}$Sb$_{0.46}$ T2SL with doping concentration $5 \times 10^{14}$ cm$^{-3}$ (after [114]).
The next two figures (Figures 30 and 31) summarize carrier lifetimes data for bulk HgCdTe and both types of T2SLs (InAs/GaSb and InAs/InAsSb) operating in MWIR (\(\lambda_{\text{cut-off}} \approx 5 \mu m\)), and LWIR (\(\lambda_{\text{cut-off}} \approx 10 \mu m\)). The trend lines HgCdTe carrier lifetimes are given after Kinch et al. [115]. InAs/InAsSb SL system shows a significantly longer minority carrier lifetime (1 \(\mu s\) for MWIR material at 77 K) in comparison to the InAs/GaSb SL operating at the same wavelength range and temperature (~100 ns).

![Figure 30. Minority carrier lifetimes versus doping concentration for MWIR HgCdTe and T2SLs at 77 K. Theoretical trend lines for n-type and p-type HgCdTe ternary alloys are taken from [115]. The dashed line for Ga-free T2SLs follows experimental data.]

![Figure 31. Minority carrier lifetimes versus doping concentration for LWIR HgCdTe and T2SLs at 77 K. Theoretical trend lines for n-type and p-type HgCdTe are taken from [115]. The dashed line for T2SLs follows experimental data.]

In lightly doped n- and p-type HgCdTe, the SRH mechanism determines the carrier lifetime. Generally, values of \(\tau_{\text{SRH}}\) are approximately more than two orders of magnitude larger for HgCdTe than those reported for the III-V semiconductors with similar bandgaps. At lower temperatures and doping level below \(10^{14} \text{ cm}^{-3}\), the HgCdTe P-i-N photodiodes become depletion limited due to SRH centers having lifetimes in the range to 10 ms [116,117]. In consequence, the potential properties of room-temperature HgCdTe photodiodes operating above 3 \(\mu m\) guarantee reaching more than order of magnitude higher detectivity (above \(10^{10}\) Jones) in comparison with value predicted by Rule 07 [118], and this detectivity is limited by the background flux [119]. Up till now, long SRH lifetime of HgCdTe gives the potential to use this material system for background limited performance (BLIP) at room-temperature operation.
5. InAsSb Bulk Photodiodes

All epitaxial techniques require large area, single crystal, and lattice-matched substrates. Ternary and quaternary layers lattice matched to InAs or GaSb substrates could be tailored to detect wavelengths in the range of 0.8 to 5 µm. As shown in Figure 32, the Ga_xIn_{1-x}As_ySb_{1-y} energy bandgap can be continuously tuned from ~475 to 730 meV while maintaining lattice-matching to a GaSb substrate [50,120], which is not retained for materials in this range such as InGaAs on InP. Both ternary (GaInSb and InAsSb) and quaternary (GaInAsSb and AlGaAsSb) materials, still not available on the market, show good performance for wavelength range ≥2 µm at the research level. The emergence of new GaInSb virtual substrates has created a promising potential for development of high performance IR detectors. Several research groups have succeeded in growing GaInSb single crystals, where there is a need to underline the properties of the Ga_{0.38}In_{0.62}Sb lattice matched to InAs_{0.35}Sb_{0.65}, with Sb composition providing the lowest energy bandgap.

![Figure 32](image_url)

Figure 32. The energy bandgap of Ga_xIn_{1-x}As_ySb_{1-y} with x and y compositions chosen in the ratio (GaSb)_{1-x}(InAs_{0.91}Sb_{0.09})_x can be tuned continuously from ~475 to 730 meV while maintaining lattice-matching to a GaSb substrate.

5.1. Technology and Properties

A variety of InAsSb photodiodes configurations have been proposed during the last forty years including n-p, n-p*, p*-n and P-i-N structures in both planar and mesa configurations. In addition, the techniques used to form p-n junctions were different, starting of Zn diffusion, through Be ion implantation in LPE technique, to more advanced epitaxial techniques such as MBE and MOCVD. The last two are predominant in fabrication of modern antimonide-based IR detectors, that essentially relies on n-type material with concentrations ~10^{16} cm^{-3}. A summary of works for the fabrication of InAsSb photodiodes is given in recently published monograph [88].

Considerable step in development of InAsSb photodiodes was obtained in 1980 when nearly lattice-matched system of InAs_{1-x}Sbx (0.09 ≤ x ≤ 0.15) and GaSb were used [9]. Slight lattice mismatch up to 0.25% for the InAs_{0.86}Sb_{0.14} epitaxial layer was accommodated in terms of low etch-pit density (~10^4 cm^{-2}). Figure 33a shows a schematic structure of a back-side-illuminated (BSI) InAs_{1-x}Sbx/GaSb photodiode. The wider-gap GaSb is a transmission window for photons reaching the InAs_{1-x}Sbx active layer and where they are absorbed. Part of the photons is also absorbed in GaSb substrate, which determines the cut-on wavelength (1.7 µm at 77 K), while the active region establishes λ_{cut-off} = 4.3 µm (see Figure 33b). The InAs_{1-x}Sbx active layer was grown using LPE technique. Both undoped n-type layer and Zn doped p-type layer show the carrier concentrations of approximately 10^{16} cm^{-3}. High quality of InAs_{0.86}Sb_{0.14} photodiodes was demonstrated by a high $R_o/A$ product in excess of 10^6 Ωcm^2 at 77 K.
P-i-N heterostructure device with an unintentionally doped InAsSb active layer (π or ν; doping level \( \sim 10^{15} \) cm\(^{-3} \)) sandwiched between P and N wider-bandgap layers have been proposed by several groups to improve device performance at near-room temperature operation. Lower diffusion dark current, higher \( R_{oA} \) product, and detectivity are related to the lack of injection of minority carriers from the wide-bandgap layers. Figure 34 shows schematic architecture of the N-i-P heterostructure and the different combinations of III-V materials for an active and contact layers. For InAsSb absorber, usually p-type GaSb and n-type InAs are used for the contact layers. Depending on the configuration of contact layers, both BSI and front-side illumination (FSI) can be used. However, for the BSI photodiode, the transparency of the substrates is also crucial. In spite of the relatively low absorption coefficients of commonly used substrates (GaSb, InAs), they are thinned to small thicknesses, even less than 10 \( \mu \)m. Sometimes, the substrates are completely etched off. What is more, many fabrication processes are not possible for InAs due to its fragility. The transparency of the substrates can be improved by using a heavy n-type doping and the use of the BM effect. A strong degeneracy of the electrons in the CB occurs already for a relatively low electron concentration (>10\(^{17} \) cm\(^{-3} \)). For example, in a heavily doped n\(^{+}\)-InAs (\( N_{d} = 6 \times 10^{18} \) cm\(^{-3} \)), the BM shift makes the corresponding substrates transparent to the 3.3 \( \mu \)m [121].

![Figure 33. BSI InAs\(_{0.86}Sb_{0.14}/GaSb\) photodiode: (a) device structure and energy-band diagram of the structure, (b) spectral response at 77 K (after [9]).](image)

![Figure 34. N-i-P double heterostructure photodiode schematic cross-section. Different combinations of III-V materials for active and cladding layers are also shown (after [91]).](image)

In addition, many of the investigations were made in the Ioffe Physical-Technical Institute, see [121–124], on different antimonide-based ternary and quaternary alloys as a materials for MWIR double heterostructure (DH) photodiodes for near-room temperature operation. These DH devices with about several-\( \mu \)m thick undoped n-InAs\(_{1-x}\)Sb\(_{x}\) active layers and p-InAs\(_{1-x-y}\)Sb\(_{x}\)P\(_{y}\) 3-\( \mu \)m-thick Zn-doped cladding contact layers are LPE grown on n-type InAs(100) substrates (with \( n = 2 \times 10^{16} \) cm\(^{-3} \) for undoped or \( n^{+} = 2 \times 10^{18} \) cm\(^{-3} \) for the Sn-doped substrates). For example, Figure 35 presents the distribution of P, Zn, Sn, and Sb atoms along the growth direction of the BSI P-InAsSbP(Zn)/p-InAs\(_{0.88}Sb_{0.12}\)(Zn)/n-InAs\(_{0.88}Sb_{0.12}/n^{+}-\)InAs(Sn) DH photodiode obtained by the secondary ion mass spectroscopy (SIMS). InAs\(_{0.88}Sb_{0.12}\) absorber with smooth Zn distribution in the vicinity of the p-n junction is optimized for the MWIR range.
Figure 35. Distribution of P, Zn, Sn, and Sb atoms along the growth direction (SIMS data) of LPE-grown P-InAsSbP(Zn)/p-InAs0.88Sb0.12 (Zn)/n-InAs0.88Sb0.12/n+-InAs(Sn) DH photodiode. Arrows on the right and left denote photon flux with respect to heterostructure layers at the photoluminescence (PL), photodiode (PD), and electroluminescence (EL) modes (after [125]).

Figure 36 shows the detector dimensional dependence of dark currents for the InAsSbP/InAs0.7Sb0.3 DH photodiodes fabricated in Ioffe Institute with active 3-μm thick n-type InAsSb region. High circular mesas were defined by a standard optical photolithography and a wet chemical etching process. Next, circular Au- or Ag-based reflective anode and cathode contacts were formed by sputtering and thermal evaporation in vacuum. Finally, flip-chip bonding/packaging procedure has been implemented. Some chips were equipped with aplanatic hyperhemispherical Si immersion lenses (Ø = 3.5 mm) with antireflection coating. After [126], InAsSbP/InAs0.7Sb0.3 DH photodiodes demonstrate diffusion-limited current at near room temperature and tunnel current at 212–250 K with predominantly series resistance in the temperature range of 270–300 K. At a room-temperature, the surface leakage current dominates the bulk one starting at mesa diameter less than ≈17 μm. As is marked in Section 3.3, the surface of semiconductor like InAs (also bulk InAsSb and InAs/InAsSb T2SLs) exhibits n-type conductivity even when the material is p-doped. After etching, the detector pixels contain exposed side-walls with a p-n+ surface junction when absorber is doped on a p-type. Thus, for good performance, such detectors would require passivation.

Figure 36. Total dark current at a reverse bias of 0.1 V (●) and 0.2 V (○) in InAsSbP/InAs0.7Sb0.3 DH photodiode at room temperature. The lines present the simulated bulk ($I_{\text{Bulk}}$) and surface ($I_{\text{Surf}}$) leakage currents as well the sum of the above two currents at a reverse bias of 0.1 V (after [126]).

Figure 37 summarizes experimental data of $R_o A$ product versus photon energy for n+-InAs/n-InAsSb/p-InAsSbP DH photodiodes in wide wavelength spectral range to 9 μm. The experimental data for InAsSb photodiodes are comparable with those of HgCdTe photodiodes produced by VIGO System. It is assumed that the photon energy in the region of 90% photocurrent drop is being close to the energy gap of photodiode active region. An exponential dependence of $R_o A$
product, approximated by \( \exp(h\nu_0/kT) \), shows that the transport properties are determined by the diffusion current and the leakage current flow mechanism is negligible.

The final construction of immersion photodiode is similar to that shown in Figure 38a [123]. Figure 38b shows the Johnson-limited detectivity at different temperatures of InAsSb DH photodiodes with immersion lenses. As seen from this figure, on the spectral responsivity characteristics, four distinct regions can be distinguished: region of a rapid responsivity decay in the wavelength range from 4.7 \( \mu \)m to 5.5 \( \mu \)m, so called the cut-off region (1), highest responsivity region with a sharp longwave response decline (2), smooth response decline region (3), and finally fast shortwave response decline region, so called cut-on region (4). A short-wave responsivity limit depends on the transparency of the substrate. In the case of a highly doped n+-InAs substrate, the BM effect shifts the detector response by 1 \( \mu \)m towards the shorter wavelengths (see difference in short wavelength spectral responsivity between heavily doped-#878 sample and undoped-#877 sample).

Figure 38. DH InAsSb immersion photodiodes: (a) construction of the immersion photodiode (after [121]) and (b) detectivity spectra at different temperatures for InAsSb photodiodes with n-InAs and n+-InAs substrates (after [123]).

Figure 39a summarizes the peak detectivity of photodiodes in dependence of temperature. The next figure, Figure 39b, presents detectivities in dependence on peak wavelength for the flat BSI photodiodes and the immersion lens photodiodes (immersion illuminated—IL). For the immersion lens photodiodes, the peak detectivities are generally about one order of magnitude higher than those for bare chip PDs—see Figure 39b. The photodiodes developed at the Ioffe Institute are superior to the majority published in literature. At the same time, the \( R_oA \) products are lower than that given in [8].
Figure 39. Detectivity of InAsSb DH photodiodes: (a) maximum detectivity and peak wavelength versus temperature (after [123]); (b) peak detectivity versus peak wavelength of photodiodes without (BSI) and with Si lenses (IL) (after [121]).

More recently, the Ioffe’s research group has demonstrated the InAsSbP/InAs$_{0.7}$Sb$_{0.3}$ DH photodiodes grown onto InAs substrates with peak responsivity about 6.5 µm. Their spectral current responsivity in temperature range between 233 K and 330 K is shown in Figure 40a. The Jonson noise limited detectivity at 233 K, achievable by a 2-stage thermoelectric cooler, is $3.2 \times 10^9$ cmHz$^{1/2}$/W at $\lambda = 6.5$ µm. The use of the integrated with photodiode a hyperhemispherical immersion Si lens ($\varnothing = 3.5$ mm) increases the $D^*$ value to $4.3 \times 10^9$ cmHz$^{1/2}$/W (see Figure 40b).

![Figure 40](image)

Figure 40. DH InAsSbP/InAs$_{0.7}$Sb$_{0.3}$ p-n photodiodes grown onto InAs substrates: (a) photoresponse spectra in the 233–330 K interval, (b) maximum detectivity for bare chip (□) and immersion lens (△) photodiodes (left scale) and responsivity (right scale,) versus temperature in a 195-µm wide bare chip (after [125]).

5.2. Performance Limits

Despite the promise of the III-V based detectors, HgCdTe remains the highest-performing IR material technology for a number of applications. High-quality InAsSb photodiodes optimized for the 3–5 µm spectral region with performance comparable to HgCdTe have been developed over the last 30 years. However, a long-standing hope that the InAsSb might become a useful material in 8–12 µm spectral band has still not been realized. In comparison with HgCdTe, the main obstacles to achieving this goal are: poor quality crystal structure (there is no ideal III–V substrate/epitaxial combination that is appropriate for LWIR spectral band), poor SRH lifetimes, and relatively high background carrier concentration (lowest at the level of $5 \times 10^{13}$ cm$^{-3}$). Moreover, the observed energy gaps of InAs$_{1-x}$Sb$_x$...
with middle Sb compositions (close to minimum energy gap (see Figures 4 and 5)) are not inherent and well controlled due to CuPt-type ordering and residual strain effects. The design of buffer layer and its thickness have strong impact on the $R_oA$ product and spectral responsivity [126].

The $R_oA$ product is a commonly encountered figure of merit for a photovoltaic detector and is defined as

$$R_oA = \left( \frac{\partial J}{\partial V} \right)^{-1}_{V_b=0},$$

at zero-biased voltage $V_b = 0$. Here $J = I/A$ is the current density, and $A$ is the detector area.

Rogalski et al. have reported a theoretical analysis of MWIR InAs$_{1-x}$Sb$_x$ ($0 \leq x \leq 0.4$) photodiodes with operation extending to the temperature range of 200–300 K [128]. Figure 41 presents a theoretical critical value of the $R_oA$ product for InAsSb DH photodiodes operating at 200 and 300 K and in spectral range between 3 and 8 μm. With the current state of technology, the electron concentration in the undoped absorber is at the level of $10^{15}$–$10^{16}$ cm$^{-3}$, thus the theoretical calculations have been done for two doping close to the level available in practice. For comparison with theoretical predictions, only limited experimental data published to 1996 is marked. The agreement is satisfactory assuming 15-μm thick base region of photodiode (solid lines). However, as is shown in this figure, the room-temperature experimental results published recently in [127], are located above theoretically predicted line. The main reason of this discrepancy is considerably lower thickness of active region—typically 4 μm. By reducing the photodiode volume in which diffusion current is generated, the corresponding $R_oA$ product increases by a factor of $L_d/d$ [3], where $L_d$ is the minority diffusion length and $d$ is the thickness of base region.

More recently, Wróbel et al. have considered the effects of doping profiles on room temperature MWIR InAsSb photodiode parameters ($R_oA$ product and detectivity) [91]. In theoretical estimates a new dependence of the spin-orbit-splitting energy on the Sb molar composition was taken into account [89] -see Figure 17. Figure 42 shows the dependence of the $R_oA$ product on the $\lambda_{cut-off}$ for InAs$_{1-x}$Sb$_x$ p-on-n and n-on-p photodiodes at 300 K. Electron and hole concentration in a 5-μm thick active regions, respectively in p-on-n and n-on-p photodiode, was assumed at the level of $10^{16}$ cm$^{-3}$. It is clearly shown that in the photodiode with p-type active region with a Sb composition close to InAs ($0 \leq x \leq 0.15$; $\lambda_{cut-off} < 4.5$ μm), Auger S mechanism considerably decreases $R_oA$ product in

Figure 41. The $R_oA$ product versus $\lambda_{cut-off}$ for InAs$_{1-x}$Sb$_x$ DH photodiodes at temperatures 200 and 300 K. The solid curves are calculated for two doping concentrations ($10^{15}$ cm$^{-3}$ and $10^{16}$ cm$^{-3}$) in the active region of photodiode with a thickness of 15 μm; the carrier concentration of $10^{18}$ cm$^{-3}$ in p$^+$-cap layer with thickness equal 1 μm is assumed. The experimental data are taken from [127] (■), [129] (○), [130] (●), [131] (□), and [132].

Figure 42. The electron and hole concentration in the undoped absorber is at the level of $10^{15}$–$10^{16}$ cm$^{-3}$. With the current state of technology, the electron concentration in the undoped absorber is at the level of $10^{15}$–$10^{16}$ cm$^{-3}$, thus the theoretical calculations have been done for two doping close to the level available in practice. For comparison with theoretical predictions, only limited experimental data published to 1996 is marked. The agreement is satisfactory assuming 15-μm thick base region of photodiode (solid lines). However, as is shown in this figure, the room-temperature experimental results published recently in [127], are located above theoretically predicted line. The main reason of this discrepancy is considerably lower thickness of active region—typically 4 μm. By reducing the photodiode volume in which diffusion current is generated, the corresponding $R_oA$ product increases by a factor of $L_d/d$ [3], where $L_d$ is the minority diffusion length and $d$ is the thickness of base region.
comparison with the photodiode with n-type absorber, in which Auger S mechanism is not significant. However, for longer wavelengths ($x \geq 0.15; \lambda_{\text{cut-off}} > 4.5 \mu m$), the n-on-p photodiode is more optimal than p-on-n one.

![Figure 42](image_url)  
**Figure 42.** The $R_o A$ product versus $\lambda_{\text{cut-off}}$ for InAs$_{1-x}$Sb$_x$ p-on-n and n-on-p photodiodes at room temperature. The theoretical lines are calculated for doping concentration of $10^{16}$ cm$^{-3}$ in both p- and n-type 5-µm thick absorber. The gathered experimental data concerns p-on-n photodiodes with n-type active region (after [91]).

The theoretical relations presented in Figure 42 were also compared with the experimental values for the p-on-n photodiode. A good agreement between both types of data was obtained, even though some experimental points are located above the theoretical line. This can probably be observed if the thickness of the active region is less than the diffusion length of the minority carriers.

Knowing the $R_o A$ product, the normalized detectivity limited by the thermal noise can be calculated using formula

$$D^* = \eta \frac{\lambda a}{2h c} \left(\frac{R_o A}{kT} \right)^{1/2}, \quad (13)$$

assuming $QE$, $\eta = 0.7$.

The next figure (Figure 43) shows experimental and theoretical normalized detectivity of p-on-n InAsSb photodiodes operating at room-temperature and different $\lambda_{\text{cut-off}}$. The discrepancy between experimental data and theoretical prediction increases versus $\lambda_{\text{cut-off}}$. It may be caused by decrease of measured $QE$, while the theoretical was assumed to be constant.

![Figure 43](image_url)  
**Figure 43.** Normalized detectivity versus $\lambda_{\text{cut-off}}$ for InAs$_{1-x}$Sb$_x$ p-on-n and n-on-p photodiodes at room temperature. The theoretical line is calculated for 5-µm thick active region with doping concentration of $10^{16}$ cm$^{-3}$. The gathered experimental data is taken from literature (after [91, 121, 133–136]).
Figure 44 compares the experimental detectivity values of commercially available different types of InAsSb photodetectors (mainly photodiodes) operating at room temperature with experimentally measured HgCdTe photodiodes and theoretically predicted curves for P-i-N higher operating temperature (HOT) HgCdTe photodiodes. All experimental data gathered in this figure indicates on the sub-BLIP photodetectors’ performance. As is shown, the best quality HOT photodiodes are fabricated by Teledyne Judson Technologies with detectivity about one order below BLIP curve for \(\lambda_{\text{cut-off}} \sim 5 \, \mu m\).

At present stage of the HgCdTe technology, the “Rule 07” metric (specified in 2007 [118]) is not a proper approach for prediction of the HgCdTe detector and system performance and as a reference benchmark for alternative technologies. The Rule 07 coincides well with theoretically predicted curve for Auger-suppressed p-on-n photodiode with electron concentration in active region ~10^14 cm^{-3} and the thickness of active region = 5 \mu m (\(t_{\text{dep}}\)-depletion thickness, \(t_{\text{diff}}\)-diffusion thickness) [119].

For sufficiently long SRH carrier lifetime in HgCdTe, which is experimentally supported at doping level below 5 \times 10^{13} \, \text{cm}^{-3}, the internal P-i-N HgCdTe photodiode current is suppressed and performance is limited by the background radiation [117]. In Extended Abstracts of The 2019 U.S. Workshop on the Physics and Chemistry of II-VI Materials [132], it was suggested to replace “Rule 07” with “Law 19”. The “Law 19” corresponds exactly with the BLIP curve for room temperature [117]. The internal photodiode current can be several orders of magnitude below “Rule 07” versus \(\lambda_{\text{cut-off}}\) and operating temperature. In this context, the alternative technologies should be considered and evaluated.

Figure 44 shows that the potential properties of HOT HgCdTe photodiodes operating in longer wavelength IR range (above 3 \mu m) guarantee achieving more than order of magnitude higher detectivity (above 10^{10} \text{Jones}) in comparison with value predicted by “Rule 07” [119]. The above estimates provide further mobilization for achieving high performance MWIR and LWIR HgCdTe FPAs operating in HOT conditions, at low production costs. In the most optimistic scenario, antimonide/arsenide-based photodetectors can totally supplant the Hg_{1-x}Cd_{x}Te devices for each spectral range and operating temperature. Particular fabrication efforts provide Hamamatsu with a variety of IR detectors with different spectra response characteristics—see Figure 45.
was placed between a bandgap gap n-type absorber region and the same narrow bandgap contact (see Figure 46a), which is key to maintain a low, di 

The p-n junction (space charge area) (see Figure 46c) is replaced by the barrier, and p contact is replaced by the n-type layer. A.M. White in his prescient patent also proposed a bias-selectable two-color detector realized by the n-type layer. Due to the absence of majority carrier flow, in this respect it is similar to a photoconductor. Its design was similar to the currently used nBn detector: a thin wide bandgap layer for device biasing. Barrier should be located near the minority carrier collector and away from the narrow-bandgap semiconductor on the other side of the barrier plays a role of a contact layer with a thickness comparable to the absorption length of light in the device, typically several microns. The narrow-bandgap semiconductor on the other side of the barrier plays a role of a contact layer for device biasing. Barrier should be located near the minority carrier collector and away from the region of optical absorption. As mentioned, the barrier needs to be carefully engineered. A nearly zero VBO should allow photogenerated holes to flow to the contact (cathode), while large offset in the CB should block majority carrier dark current, re-injected photocurrent.

The nBn detector can be considered as a hybrid of the p-n photodiode and the photoconductor. The p-n junction (space charge area) (see Figure 46c) is replaced by the barrier, and p contact is replaced by the n-type layer. Due to the absence of majority carrier flow, in this respect it is similar to a photoconductor with unity gain.

The operating principles of the nBn and related detectors have been widely demonstrated in the literature [101,142–150]. The idea of nBn concept has originated with bulk InAs [142]. However, it is difficult to realize little or no VBO using this material and other bulk IR detector materials such as InSb and HgCdTe. In the middle of first decade of the XXI century, the IR technology has changed after the introduction of 6.1 Å III-V material family and first demonstration of high-performance detectors and FPAs [142,143]. The introduction of broken-gap T2SLs was particularly helpful in the design of unipolar barriers, as they enable any modification of the detector architecture with better control of unipolar barriers, as they enable any modification of the detector architecture with better control of
band edge alignments [151]. The ability to tune the positions of the CB and VB edges independently is easier than in bulk materials.

Figure 46. Bandgap diagram of nBn (a) and nBp (b) barrier detectors (Δ-VBO) and p-n photodiode (c).

In general, unipolar barriers architecture is used to increase the photogenerated carriers’ collection efficiency, reduce dark current (associated with SRH processes), and noise without inhibiting photocurrent flow. In particular, the barrier plays a role of self-passivation and serves to reduce the surface leakage current. Other key benefits of uniform doping of nBn detector (n-type barrier, Bn) is the absence of depletion regions in the narrow-gap absorption layer. It offers a way for materials with relatively poor SRH lifetimes, such as all III-V compounds, to grow on lattice-mismatched substrates such as GaAs, with a reduced penalty of excess of the dark current generated by SRH centers associated with dislocations.

The papers [144,145] established a criterion for combination of voltage and barrier doping concentration allowing operation with no depletion region in the narrow gap active layer. In some material system, a barrier in the VB is present when the barrier layer is n-type (Bn), that can significantly impede hole current flow from absorber to the contact layer. In such situation, a relatively high voltage (so called “turn-on voltage”) is required to overcome. A p-type doping of the barrier, Bp, helps to reduce the barrier in the VB (see Figure 46b) but rather a potential well for holes in the VB which does not impede hole transport between the absorption layer and the contact layer. However, in the last case, a p-type barrier inherently causes depletion regions to form in the narrow-gap absorption layer for all voltages, what should be avoided (depletion regions cause excessive G-R dark currents).

Detailed growth procedures and device characterization of MWIR detectors based on InAsSb/AlAsSb material system were published in several papers, e.g., References [144,145,147,152,153]. The principal layers of the device structures were n-type doped and consisted of a thick InAs1−xSbx absorption layer (1.5–3 μm), a thin AlAs1−xSbx barrier layer (0.2–0.35 μm), and a thin (0.2–0.3 μm) InAs1−xSbx contact layer. The bottom contact layer was highly doped. InAsSb/AlAsSb nBn structures were grown on either GaAs(100) or GaSb(100) substrates by using MBE technique [153]. The n-type doping was usually obtained by either Si or Te elements. The remaining structures were grown directly onto lattice-matched GaSb (100) substrates, while lattice-mismatched GaAs (100) substrates were used, structures were grown on a 4-μm thick GaSb buffer layer.

Figure 47a shows an example of such an nBn structure that was theoretically investigated by Martyniuk and Rogalski [154]. The alloy compositions of x = 0.09 to 0.2 for the InAs1−xSbx absorber layers provided the λcut-off wavelengths of ~4.1 to ~5.0 μm at 150 K, respectively. The current-voltage characteristics were taken from [155]. The dark current density for nBn InAs0.80Sb0.195 structure was 1.0 × 10−3 A/cm² at 200 K and 3.0 × 10−6 A/cm² at 150 K.
where $N$ varies as $1/N$. Sensors 2020 × concentration 5

Kinch has modeled the diode characteristics for InAsSb close to InAs. SRH mechanisms can be calculated as for InAsSb/AlAsSb nBn structures were grown on either GaAs(100) or GaSb(100) substrates by using a thin (0.2–0.3 μm) InAs1–xSbx absorption layer (1.5–3 μm at 150 K) tied together in parallel (after [155]), and a potential well for holes is no depletion region. In the low-temperature region, for temperatures below $T_c$, the nBn detector exhibits a higher signal-to-noise ratio in comparison with the conventional p-n diode operating at the same temperature, or otherwise the nBn detector can operate at a higher temperature than a conventional p-n diode with the same value of dark current. InAsSb p-n photodiode is characterized by a higher crossover temperature, $T_c$, between diffusion and G-R contribution than InSb, which results from the larger bandgap of InAsSb than InSb (only in a certain range of Sb molar composition, for InAsSb close to InAs).

In nBn InAsSb detector, the depletion current is totally suppressed (see Figure 47c) because there is no depletion region. In the low-temperature region, for temperatures below $T_c$, the nBn detector exhibits a higher signal-to-noise ratio in comparison with the conventional p-n diode operating at the same temperature, or otherwise the nBn detector can operate at a higher temperature than a conventional p-n diode with the same value of dark current. InAsSb p-n photodiode is characterized by a higher crossover temperature, $T_c$, between diffusion and G-R contribution than InSb, which results from the larger bandgap of InAsSb than InSb (only in a certain range of Sb molar composition, for InAsSb close to InAs).

In InAsSb nBn detector (n-type absorber), the dark current density limited by the Auger 1 and SRH mechanisms can be calculated as

$$J_{\text{dark}} = \frac{qN_d^2}{2t^2} + \frac{q\mu_i^2 t}{2t} \tau_{\text{SRH}},$$

where $N_d$ and $t$ are the doping and thickness of active region, respectively. The influence of G-R current and of depletion region (from $j_{\text{dark}} = q\mu_i^2 t\tau_{\text{SRH}}$, were $w$ is the depletion width) can be omitted.

According to Equation (14), the Auger 1 generation varies as $N_d$, whereas the SRH generation varies as $1/N_d$. It follows that that the diffusion dark current mostly depends on absorber doping concentration and on value of SRH lifetime. Kinch has modeled the diffusion current in an InAsSb absorber versus doping concentration at 150 K (assuming $\tau_{\text{SRH}} = 400$ ns (see Figure 19) and an absorber thickness of 3 μm). As is shown in Figure 48a, the optimum doping is $10^{16}$ cm$^{-3}$. Next, Figure 48b shows the dark current density for InAs$_{0.91}$Sb$_{0.09}$ absorber versus inverse temperature for doping concentration $5 \times 10^{15}$ cm$^{-3}$. This figure shows that the dark current is equal to the $f/3$ background.
flux at temperature of ~175 K. At 150 K, the dark current density is $2 \times 10^{-7}$ A/cm$^2$, in agreement with experimental nBn InAsSb data [156]. In this way, the nBn InAsSb detector operating with f/3 optics reaches BLIP for $T > 160$ K (temperature two times higher than traditional InSb).

**Figure 48.** Dark current density for InAsSb nBn absorber ($\tau_{SRH} = 400$ ns, $t = 3$ $\mu$m): (a) versus doping concentration, and (b) versus inverse temperature for a donor doping $5 \times 10^{15}$ cm$^{-3}$ (after [78]).

Interesting difference in dark current density versus temperature is shown in Figure 49 for two nominally identical nBn devices, each operating at a bias of $-0.1$ V. The devices differ only in the doping of the barrier layer. The nB$_n$ device with a uniform n-type doping through the entire structure exhibits a single straight line, typical for the diffusion limited behavior. The nB$_p$ device, with a barrier doped opposite to the absorber, exhibits two-slope behavior, typical for the diffusion limited mode at high temperatures and the G-R limited behavior at low temperatures. As is shown, the dark current density at 150 K is more than two orders of magnitude greater for the device with the p-type due to the presence of the depletion region. It appears that for a typical QE of 70% at f/3 optics, the BLIP temperature is about 140 K compared with ~175 K for the detector with the n-type barrier.

**Figure 49.** Dark current density vs. temperature for two identical InAsSb/AlSbAs nBn devices with opposite barrier doping. Active layer bandgap wavelength is 4.1 $\mu$m at 150 K (after [150]).

Over present-day IR technologies, InAsSb-based barrier detectors offer numerous advantages, such as reduced depletion and surface leakage currents, normal-incidence absorption. However, the promise of the excellent performance of these detectors has yet to be practically realized.
Martyniuk and Rogalski have predicted the temperature dependencies of dark currents of different types of IR detectors, including InAsSb nBn detectors, optimized at a $\lambda_{\text{cut-off}} = 5 \, \mu m$ [157].

The advantages of the nBn structure, compared to the InAsSb photodiode with the built-in depletion region, are clearly visible. The nBn structure enables operation at much higher temperatures while maintaining the same dark current values (see Figure 50). However, still these values are not comparable to the HgCdTe photodiodes with p-on-n or even n-on-p design. More comparable to HgCdTe photodiodes is T2SL InAs/InAsSb nBn detector. MWIR HgCdTe systems operating at f/3 optics at 160 K are commercially available.

![Figure 50. Dark current density versus temperature for InAsSb p-on-n photodiode and nBn detector, T2SL nBn detectors, and HgCdTe photodiodes with $\lambda_{\text{cut-off}} = 5 \, \mu m$.](image)

Early efforts in development of InAsSb barriers detectors focused on lattice-matched structures to GaSb substrates with $\lambda_{\text{cut-off}} \sim 4.1 \, \mu m$. Different strategies were attempted to extend the $\lambda_{\text{cut-off}}$ above 5 $\mu m$ including various metamorphic buffer layers and InSb interface ML into the InAsSb [158,159]. It appears that the activation energies estimated from temperature dependence of dark currents is higher than the bandgap of InAsSb absorbers, which is explained by location of the accumulation layer between the barrier and absorber due to band bending [160]. Some barrier detectors also exhibits an increase in the turn-on bias versus temperature as a result of the Fermi level position changes in low-doped or unintentionally doped absorbers. At higher temperature, a higher voltage is needed to deplete the barrier-absorber interface region [161].

Lin et al. [42] have described bulk InAsSb barrier detectors with a $\lambda_{\text{cut-off}}$ about 10 $\mu m$ at 77 K. The estimated InAs$_{0.60}$Sb$_{0.40}$ absorber parameters at 77 K are as follows: the hole mobility of $10^{3} \, cm^{2}/Vs$, the minority hole lifetime of 185 ns, and the diffusion length of 9 $\mu m$. The current-voltage characteristics have been influenced by G-R component associated with a depletion region adjacent to the barrier, as well as probably tunneling components. To reach diffusion-limited dark current, the VBO associated with the heterointerfaces must be eliminated.

The spectral detectivity curves of LWIR nBn InAsSb detector, with two absorber’s doping levels, are shown in Figure 51. The demonstrated detectivity of $2 \times 10^{11} \, cmHz^{1/2}/W$ at 2\pi FOV and wavelength of 8 $\mu m$ was estimated in spite of the significant shift towards shorter wavelengths (BM shift) with doping. For 1 $\mu m$ thick InAs$_{0.60}$Sb$_{0.40}$ absorber at $\lambda = 8 \, \mu m$, an absorption coefficient was estimated at $3 \times 10^{3} \, cm^{-1}$, what implies QE = 22%. The QE increases with bias, until it reaches a constant level for $-0.4 \, V$ and with thickness of absorption layer (to 40% for 3-$\mu m$ thick absorber). Generally, due to poor quality crystal structure, the performance of LWIR bulk InAsSb detectors are considerably inferior in comparison with HgCdTe photodiodes.
The LWIR InAs T2SLs is easier compared to the InAs Sensors achieved at about 130 K and is close to HgCdTe photodiode. The BLIP performance with temperature for diode implementation in the MWIR range, where they could have an advantage over the InAs dark currents. In addition, the depletion region in the surface p-n junction is a source of both G-R and tunneling + diffusion current. For InAsSb SL detectors, it is better to use an n-type absorber due to their surface properties. In the last decade, a rapid progress in their fabrication is made due to some defect tolerance and its robust material properties. In general, the growth of the InAs/InAsSb T2SLs is easier compared to the InAs/GaSb ones and InAs/InAsSb SLs have longer minority carrier lifetimes (see Section 4.3). On the other hand, its spectral tuneability is somewhat smaller and the InAs/InAsSb T2SLs has weaker optical absorption. In addition, vertical hole transport properties, especially in the LWIR range, are more challenging.

### 6. InAs/InAsSb Superlattice Detectors

Development of the T2SLs has more than forty years history. Figure 52 gives it a short description in the form of a timeline, including InAs/InAsSb SLs. The development of InAs/InAsSb SLs for IR detector applications is described in detail in several papers [19,21,159,162]. It has a long and interesting history that predates the InAs/GaSb SL detectors. Up till 2010 this material system was much less explored than the T2SLs InAs/GaSb. In the last decade, a rapid progress in their fabrication is made due to some defect tolerance and its robust material properties. In general, the growth of the InAs/InAsSb T2SLs is easier compared to the InAs/GaSb ones and InAs/InAsSb SLs have longer minority carrier lifetimes (see Section 4.3). On the other hand, its spectral tuneability is somewhat smaller and the InAs/InAsSb T2SLs has weaker optical absorption. In addition, vertical hole transport properties, especially in the LWIR range, are more challenging.

#### 6.1. Performance Limits

The InAs/GaSb SL detectors are most commonly designed with p-type absorbers, while in the case of InAs/InAsSb SL detectors, it is better to use an n-type absorber due to their surface properties. In InAs/InAsSb SLs, the surface Fermi level is pinned above the CB minimum, resulting in a degenerate n-type surface. Thus, for the n-type absorber no special surface treatment is required since the surface band-bending potential repels the minority carriers (holes) away from the surface. In a p-type absorber, the minority electrons are attracted to the n-type surface contributing to the surface leakage current. In addition, the depletion region in the surface p-n+ junction is a source of both G-R and tunneling dark currents.

Summarizing the above discussion—choosing between the two types of SLs, InAs/GaSb and InAs/InAsSb, it is necessary to take into account the λ_{cut-off}, the ease of growth technology and processing capabilities (including passivation). In general, the InAs/InAsSb SLs are more effective for implementation in the MWIR range, where they could have an advantage over the InAs/GaSb ones.

Figures 50 and 53 compare the predicted dependence of dark current density on operating temperature for different material systems including InAs/InAsSb SLs with λ_{cut-off} of 5 and 10 μm. The LWIR InAs/GaSb T2SL materials have larger dark current densities than HgCdTe. Their inferior parameters appear to result from a relatively high background concentrations (about 10^{15} cm^{-3}) and a shorter minority carrier lifetime. The BLIP performance with f/1 optics for InAs/InAsSb nBn detector is achieved at about 130 K and is close to HgCdTe photodiode.
leakage current. In addition, the depletion region in the surface p-n+ junction is a source of both G-R temperature for different material systems including InAs/InAsSb SLs with implementation in the MWIR range, where they could have an advantage over the InAs/GaSb ones.

Generally, however, the InAs/InAsSb SL detectors are characterized by lower QE ~ 40–50% (see Figure 55) than HgCdTe photodiodes (>70%), mainly due to lower absorption coefficient. For good photoresponse of the nBn detectors, the devices must be biased in a shorter minority carrier lifetime. The BLIP performance with f/1 optics for InAs/InAsSb SL detectors, it is better to use an n-type absorber due to their surface properties.

In general, the InAs/InAsSb SLs are more effective for processing capabilities (including passivation). In general, the InAs/InAsSb SLs are more effective for implementing in the MWIR range, where they could have an advantage over the InAs/GaSb ones.

Figure 52. Roadmap development of T2SL IR photodetectors including InAs/InAsSb SL photodiodes and barrier detectors.

Figure 53. Dark current density versus temperature for T2SL nBn detectors and HgCdTe photodiodes with a $\lambda_{\text{cut-off}} = 10 \mu m$.

Figure 54 compares the $R_o A$ and $R_d A$ values for InAs/InAsSb SL and HgCdTe devices operating in a wide IR spectral range. For good photosresponse of the nBn detectors, the devices must be biased typically in the range above ~100 mV. Thus, the $R_d A$ product is used instead of the $R_o A$, where $R_d$ is the...
dynamic resistance at non-zero bias voltage. The solid line is the trend line calculated with Teledyne empirical model (“Rule 07” [118]) for P-on-n HgCdTe structures. The HgCdTe experimental data are given for photodiodes fabricated by Raytheon Vision Systems (RVS). As we can see, the upper results for InAs/InAsSb SL detectors rival that of practical HgCdTe devices.

Figure 54. $R_c A$ and $R_d A$ product of nonbarrier and barrier T2SL InAs/InAsSb detectors on $\lambda_{\text{cut-off}}$ wavelength dependence compared to experimental data and trend line (“Rule 07”) for HgCdTe photodiodes at 77 K. The experimental data for InAs/InAsSb SLS devices are taken from [163–168].

In the case of InAs/InAsSb SL barrier detectors having a $\lambda_{\text{cut-off}} \sim 12 \mu m$, the values of $R_c A > 10^7 \text{ cm}^2$ at 77 K are measured. These values are higher than those measured for HgCdTe photodiodes. Generally, however, the InAs/InAsSb SL detectors are characterized by lower QE ~ 40–50% (see Figure 55) than HgCdTe photodiodes (>70%), mainly due to lower absorption coefficient.

Figure 55. QE versus thickness of active region for InAs/InAs$_{1-x}$Sb$_x$ T2SL photodetectors at 77 K. The theoretical prediction (solid line) is compared with the experimental QE reported by different research groups as indicated inside the figure (adapted after [169]).

The main technological challenge for the fabrication of SL-based devices is the growth of high-quality active region with sufficient thickness to achieve satisfactory QE. For InAs/InAsSb photodetectors, a 5 $\mu$m thick absorber is sufficient to provide high enough QE. It may provide QE about 50%, due to a longer effective lifetime and diffusion length, in comparison with InAs/GaSb SLs. Figure 54 shows the theoretical QE of InAs/InAsSb T2SL detectors versus the absorber thickness, calculated for the effective carrier lifetime of 150 ns [169]. Table 4 summarizes reports of InAs/InAsSb-based SL IR detector results.
Table 4. InAs/InAsSb superlattice infrared detectors.

|                  | Sl. Period [nm] | Polarity | Size [µm] | Absorber Thickness [µm] | T [K] | $I_0 (T, V)$ [A/cm²] | $R_v$ (A/T,V)×QE [A/W] | $D^* (A, T, V, η)$ [cm²s/Hz¹/₂/W] | References |
|------------------|----------------|----------|-----------|--------------------------|-------|-----------------------|--------------------------|---------------------------------|------------|
| InAs/InAsSb$_{0.35}$ | 18.4           | nBn      | s = 410   | 2.2                      | 77    | $5 \times 10^{-6}$    | $0.24 (12, 77, -0.3)$     | $1 \times 10^6 (12, 77, -0.3)$   | Kim et al. [163] |
| InAs/InAsSb$_{0.43}$ | 14.7           | d = 100–400 | 2.3       | 77                       | 14.6  | $0.7 (77, -0.3)$      | $4.8 (12.8, 77, -0.3)$     | $4.6 \times 10^{11} (4.5, 150, -0.2)$ | Hoang et al. [164] |
| Dual band        | 10.5           | pN       | s = 100–400 | 2.6                      | 77    | $4.4 \times 10^{-6}$  | $3.5 (7.9, 77, -0.09)$     | $2.8 \times 10^{11} (7.9, 77, -0.09)$ | Haddadi et al. [165] |
| InAs/AlAsSb/InAs/ InAs$_{0.45}$Sb$_{0.55}$ | 6.3/10.5       | nBn      | s = 100–400 | 2                        | 77    | $5.1 \times 10^{-7}$  | $1.5 (4, 77, 0.1)$          | $8.2 \times 10^{12} (4, 77, 0.1)$   | Haddadi et al. [166] |
| InAs/InAsSb$_{0.30}$ | 12              | nBn      | s = 100–400 | 2                        | 77    | $8 \times 10^{-5}$    | $2.65 (7.5, 77, -0.8)$     | $4.7 \times 10^{11} (7.5, 77, -0.08)$ | Haddadi et al. [167] |
| Dual band        | 22/25          | nBn      | s = 100–400 | 4                        | 77    | $1.13 \times 10^{-4}$ | $3.37 \times 10^{-4}$ (1.8, 77, 0.04) | $5.1 \times 10^{12} (6.45, 77, 0.04)$ | Haddadi et al. [168] |
| InAs/InAsSb$_{0.35}$ | 5.5             | nBn      | d = 60–300 | 3                        | 150   | $4 \times 10^{-3}$ (150, -2.5 V) | $1.35 (4.1, 208, 0.3 V)$ | $2.6 \times 10^{11} (4.1, 208, 0.3 V)$ | Teledyne Judson [159] |
| InAs/InAsSb$_{0.34}$ | 5               | nBn      | s = 250    | 2.6                      | 150   | $4.5 \times 10^{-4}$ (150, -0.2 V) | $-52$% | $4.6 \times 10^{11} (4.5, 150, -0.2 V)$ | Ting et al. [171,172] |
| InAs/InAsSb$_{0.35}$ | 12              | d = 400   | 2          | 77                       | 8.0   | $9 \times 10^{-5}$ (77, -20 mV) | $1.26 (7.5, 77, -0.21)$ | $5.4 \times 10^{10} (7.5, 77, -0.21)$ | Wu et al. [175] |
| InAs/InAs$_{0.7}$Sb$_{0.3}$ | 6.6             | d = 400   | 2          | 77                       | 150   | $3.3 \times 10^{-4}$ (150, -20 mV) | $1.76 (4, 150, 0.04)$ | $1.2 \times 10^{11} (4.5, 150, 0.02)$ | Wu et al. [174] |
| InAs/InAsSb$_{0.35}$ | 5.15            | nBn      | d = 60–310 | 3                        | 150   | $1 \times 10^{-3}$ (150, -0.5 V) | $7.1 \times 10^{11} (4.4, 150, -0.05)$ | Durlin et al. [175] |
| InAs/InAsSb$_{0.35}$ | 7.4             | pBn      | d = 100–400 | 2.1                      | 150   | $4.4 \times 10^{-5}$ (150, -0.05 V) | $1.48 (4, 150, -0.05)$ | $47$% | $7.1 \times 10^{11} (4.4, 150, -0.05)$ | Wu et al. [176] |
| InAs/InAsSb$_{0.35}$$_{0.16}$ monolithically Si-integrated | 13.3           | nBn      | d = 50      | 4                        | 200   | $1.4 \times 10^{-2}$ (200, -0.1) | $0.88 (5, 200, -0.1)$ | $1.5 \times 10^{10} (5.0, 200, -0.1)$ | Delli et al. [177] |
| InAs/InAsSb$_{0.3}$ | 13.5            | p-n      | d = 320    | 1                        | 150   | $9.1 \times 10^{-6}$ | $0.78 (4.5, 150)$ | $3.4 \times 10^{10} (4.5, 150)$ | Wu et al. [178] |
| InAs/AlAsSb/InAs/InAsSb | -               | p-i-n    | s = 100–400 | 1                        | 300   | $9.6 \times 10^{-3}$ (300, -0.05) | $0.47 (1.8, 300, 0.37)$ | $6.45 \times 10^{10} (1.8, 300, -0.05)$ | Haddadi et al. [179] |

$\text{d}$ - diameter of detector; $s$ - square detector.
Figure 56 shows the comparison of the simulated detectivity of P-on-n HgCdTe photodiodes versus $\lambda_{\text{cut-off}}$ and operating temperature with the experimental data T2SL InAs/InAsSb photodetectors operating at 78 K, 150 K and 200 K. The solid lines present theoretical thermal limited detectivity for HgCdTe photodiodes, modeled using the “Rule 07” dark current density. This figure indicates that the measured thermally limited detectivity of T2SLs InAs/InAsSb photodetectors exhibits lower performance in comparison to the HgCdTe photodiodes not even reaching the HgCdTe level. That is mainly related to the fairly high background concentrations ($\sim 1 \times 10^{15}$ cm$^{-3}$) and a lower absorption coefficient and in consequence lower QE. Improvement in those two fundamental parameters seems to be critical to reach the theoretically predicted T2SLs photodiodes performance.

![Figure 56](image_url)

**Figure 56.** The simulated detectivity of the P-on-n HgCdTe detectors (Rule 07, $QE = 70\%$) versus $\lambda_{\text{cut-off}}$ and temperature. The experimental data is taken with different sources (as marked) for InAs/InAsSb T2SL photodetectors operating at 78 K, 150 K, and 200 K.

6.2. Superlattice Detector Structures

The majority of the T2SL devices are grown by MBE being expensive for mass production, while the recent interest in T2SLs growth by MOCVD is observed which is related to the low-cost and versatile production guaranteed by that technique [173,174].

In the last decade, the particular contribution in development of T2SLs InAs/InAsSb detectors and FPAs has given two research groups, group at Jet Propulsion Laboratory (JPL) [107,158,159,161,171,172] and the Razeghi group from Northwestern University [164–168,173,174]. Both groups have demonstrated the versatility of the T2SLs InAs/InAsSb with operating $\lambda_{\text{cut-off}}$ ranging from 2 to 14 μm and beyond. In this way, SLs material expends the limited spectral range covered by traditional bulk III-V IR detectors based on InGaAs and InSb.

Table 4 indicates that the T2SLs InAs/InAsSb detectors are fabricated mainly in two configurations: p-n junction photodiodes (impurity diffusion [178], MBE [164], MOCVD [173,174]) and nBn barrier detectors (mainly MBE). The T2SL InAs/InAsSb photodiodes, as shown in Figure 34, are mainly constructed on P-i-N heterostructures. The low minority carrier concentration in the high bandgap materials allows for the suppression of diffusion dark current and reaching higher $R_dA$ product and detectivity.

In comparison with the p-n junction T2SLs InAs/InAsSb photodiodes, SLs IR barrier devices are more effective (see Figures 50 and 53). As is described in Section 5, the unipolar barrier structures allow reducing G-R dark current by suppressing SRH mechanism and surface leakage. That is particularly useful for the p-n devices based on III-V materials, inherently exhibiting excessive depletion dark current and suffering from the lack of proper surface passivation.
InAsSb-based bulk and SLs photodetector performance has been increased by the monolithically integrated microlenses based on GaAs/GaSb substrates [180,181]. The hemispherical or hyperhemispherical microlenses increase the effective detector’s optical area by $n_r^2$ of $n_s^2$, respectively, where $n_r$ is the index of refraction of the lenses [182]. The detectivity gain achieved with hyperhemispherical immersion is $n_r^2$. Since the refractive index of GaAs is 3.4, a factor of 10 improvements in the detectivity is observed for n-type and p-type InAsSb absorbers [180]. Figure 57 shows the immersion lens formed from GaAs substrates using numerically controlled micromachining.

**Figure 57.** The photodetector assembled on the 3-stage Peltier thermoelectrical cooler and GaAs immersion microlens. The improvement in detectivity reached by hyperhemispherical microlens is proportional to $n_r^2$, where $n_r$ is the GaAs refractive index (after [180]).

The nBn (pBp) detector design is also useful for the realization of dual-band detection capabilities shown schematically in Figure 58. The dual-band barrier structures could be easily fabricated due to the highly developed III-V materials growth techniques. Typical dual-band, barrier T2SLs detector consist of the two narrow bandgap absorbers optimized for the two different IR spectral regions being separated by ~0.1 μm thick wide bandgap barrier for majority carriers (no barrier for minority carriers is assumed). That architecture blocks the majority carrier current between the two electrodes by the large energy offset, while there is no barrier for photo-generated minority carriers. Both barriers should be characterized by negligibly small VB or CB discontinuity with respect to both n-type (p-type) doped active regions. The absorbers (channels) are addressed consequently by the polarity of the applied voltage. Detector with pBp design exhibits faster operation with lower integration times, due to the higher minority carrier mobility (electrons).

**Figure 58.** Schematic of nBn (a) and pBp (c) T2SL dual-band detector operations under forward and reverse bias. Schematic view of dual-band MWIR/LWIR detector architecture is shown in centre (b).
Krishna et al. have demonstrated two-color barrier detectors based on nBn or pBp InAs/GaSb SL architectures [183,184]. Furthermore, their improvement in InAs/InAsSb two-band SL fabrication was demonstrated in [166,168].

6.2.1. Photodiodes

Both MWIR [174,185] and LWIR [164,173] T2SLs InAs/InAsSb photodiodes have been demonstrated. The experimentally measured dark current density of MWIR photodiodes with a $\lambda_{\text{cut-off}} = 5.4$ μm at 77K, was found larger than T2SLs InAs/GaSb detectors [186]. This was related to the higher probability of the carrier tunneling due to reduced VBO and CBO in T2SLs InAs/InAsSb.

The better quality LWIR T2SLs InAs/InAsSb photodiodes have been demonstrated by the Razeghi group [164]. Even if the Sb composition increases ($x_{\text{Sb}} = 0.43$), the material quality is still high leading to the detector’s high performance. The absorber $\lambda_{\text{cut-off}}$ depends on the VB level in the InAsSb layer being directly related to the Sb composition. The presented samples were grown by MBE on Te-doped (001) GaSb substrate. The analyzed device is built of 0.5-μm InAsSb buffer layer, a 0.5-μm bottom Si-doped n-type contact ($N_D \sim 10^{18}$ cm$^{-3}$), a 0.5-μm thick n-type barrier, a 2.3-μm Be-doped p-type active region ($N_A \sim 10^{15}$ cm$^{-3}$), and a 0.5-μm top p-type ($N_A \sim 10^{18}$ cm$^{-3}$). Finally, the structure was capped with a 200-nm p-type GaSb layer.

Figure 59 shows the electrical characteristics of the LWIR InAs/InAsSb SL photodiode within temperature range from 25 to 77 K. At 77K the $R_oA$ product is 0.84 Ωcm$^2$ indicating lower value in comparison with InAs/GaSb counterpart. For $T > 50$ K, the diode exhibits an Arrhenius type behavior with related activation energy, $E_a = 39$ meV meeting the relation $E_a \sim E_g/2$ of the absorber (~80 meV for 15 μm) indicating on the active region G-R current as the limiting mechanism. For $T < 50$ K, the $R_oA$ product does not follow the trend line and is less sensitive versus temperature. This behavior indicates on the other mechanisms either the tunneling or surface leakage at this range of temperature.

![Figure 59](image)

Figure 59. Dark electrical characteristics of LWIR InAs/InAsSb SL photodiodes: (a) current-voltage characteristics versus temperatures; (b) $R_oA$ versus temperature (after [164]).
The spectral characteristics of photodiode are presented in Figure 60. At 77 K, the sample exhibited a 100% \( \lambda_{\text{cut-off}} = 17 \ \mu m \) and 50% \( \lambda_{\text{cut-off}} = 14.6 \ \mu m \). High QE is reached when biased >150 mV and saturates at 300 mV where the peak current responsivity reaches 4.8 A/W, corresponding to a QE of 46% for a 2.3 \( \mu m \) thick absorber. Detectivity shown in Figure 60b, limited by shot noise and Johnson noise, is estimated using equation

\[
D^* = R_i \left( 2eH_{\text{dark}} + \frac{4kT}{RA} \right)^{1/2},
\]

where \( R_i \) is the device responsivity.

The effective passivation of T2SLs InAs/InAsSb photodiodes is in a very early stage of development. Usually, photodiodes are not passivated. The simplest passivation is based on the common dielectric insulators deposited onto exposed surface of the device, utilized in the silicon industry (such as an oxide or nitride of silicon). In the case of T2SLs InAs/InAsSb photodiodes, the passivation challenges are caused by the well-known surface accumulation of InAsSb. Sulphur anodic was found to be more effective passivant for p\(^+\)B\(_n\)n\(^+\) detector structure compared to fluoride and chloride anodic films [186]. In [187] the dark current suppression was observed in 8-\( \mu m \) \( \lambda_{\text{cut-off}} \) T2SLs InAs/InAsSb structure with combination of a double-electron barrier and SiO\(_2\) passivation. In addition, the effect of hydrogenation could be taken into account for passivation of bulk defects in MWIR T2SLs InAs/InAsSb [188].

6.2.2. Barrier Detectors

Klipstein et al. [189] proposed the division of the barrier detectors into two groups: X\(_B\)n and X\(_B\)p, where X is a contact layer where doping, material, or both can be varied. Usually, however, the most popular are four types of SLs barrier detectors schematically described in Table 5.
The detector is built of lightly p-type T2SLs InAs/GaSb active layer being sandwiched between n-type T2SLs InAs/AlSb hole barrier (hB) and wider T2SLs InAs/GaSb electron barrier (eB). The barriers are designed to exhibit zero CBO and VBO in relation to the active layer. A heavily doped n-type InAsSb layer adjacent to the eB plays the role of the bottom contact. The SRH and trap-assisted tunneling (TAT) are reduced by the N-p heterojunction between the hB and active layer.

### Table 5. T2SLs barrier detectors.

| Flat-Band Energy Diagrams | Example of Detector Structure | Description | Reference |
|---------------------------|-----------------------------|-------------|-----------|
| nBn                       | ![Detector Structure](image) | The top contact, absorber, and bottom contact layers are built of T2SLs InAs/InAsSb. As a unipolar barrier located between the top contact and absorber, about 100-nm thick AlAsSb Be-doped to 10^{15} cm^{-3} is used. The top contact and absorber layers are n.i.d while the bottom contact is Te-doped to 10^{17} cm^{-3}. | [171] |
| pBn barrier               | ![Detector Structure](image) | The epitaxial growth starts with a 700 nm thick n"-doped GaSb/InAs/AlSb buffer layer, next a 500 nm thick n"-doped bottom contact layer, a 2.1 μm n.i.d active layer, a 500 nm electron barrier, and a 300 nm thick p-type top contact. The top/bottom contacts and active layer are built of T2SLs InAs/InAsSb. Si and Be are used for n-type and p-type dopants, respectively. | [176] |
| pBp                       | ![Detector Structure](image) | The contact layers and absorber are built of T2SLs InAs/GaSb. The barrier layer is based on a T2SLs InAs/AlSb. The proper lattice match to the GaSb substrate is ensured by InSb-like interfaces. | [189] |
| CBIRD (Complementary barrier infrared detector) | ![Detector Structure](image) | The detector is built of a lightly p-type T2SLs InAs/GaSb active layer being sandwiched between n-type T2SLs InAs/AlSb hole barrier (hB) and wider T2SLs InAs/GaSb electron barrier (eB). The barriers are designed to exhibit zero CBO and VBO in relation to the active layer. A heavily doped n-type InAsSb layer adjacent to the eB plays the role of the bottom contact. The SRH and trap-assisted tunneling (TAT) are reduced by the N-p heterojunction between the hB and active layer. | [190] |
Main activity in development of the T2SLs InAs/InAsSb barrier detectors is directed to nBn devices operating in MWIR spectral range at much higher temperatures than InSb photodiodes. Generally, however, these detectors can be optimized for wider spectral range up to VLWIR [191]. The unipolar hole barriers are only available for T2SLs InAs/GaSb and mostly for LWIR. In addition, there is no good hole barrier for T2SLs InAs/InAsSb. Figure 61 shows same images taken with JPL T2SLs arrays in wide IR spectral range. In this section the representative examples of barrier detector performance are presented.

![Image of T2SLs InAs/InAsSb barrier detectors](image_url)

**Figure 61.** Images from JPL antimonide bulk and T2SLs barrier FPAs with λ_{cut-off} in wide IR range (after [159]).

The JPL nBn device structure grown by MBE on GaSb substrate is similar to that shown in Figure 47a. The top/bottom contacts and active layers consist of 16/20 and 525 periods of (37Å, 13Å) T2SLs InAs/InAsSb, respectively. As a unipolar barrier located between the top contact and absorber, 120-nm thick AlAs_{0.085}Sb_{0.915} Be-doped to 10^{15} cm^{-3} is used. The top contact and active layers are n.i.d, while the bottom contact is Te-doped to 10^{17} cm^{-3}.

The experimentally measured dark current density versus applied voltage for temperature range between 89 K and 222 K is shown in Figure 62. The inset shows an Arrhenius plot of dark current under bias −0.2 V with activation energy of 0.205 eV. The dark current density at 157 K and bias voltage of −0.2 V is 9.6 × 10^{-5} A/cm², which is ~4.5 higher than “Rule 07” for λ_{cut-off} = 5.4 μm.

![Graph of dark current density vs. voltage](graph_url)

**Figure 62.** Dark current density versus voltage, for temperatures 89–222 K. The inset presents the dark current temperature dependence −0.2 V (after [171]).

The QE shown in Figure 63 is derived from measured responsivity of the BSI samples through the GaSb substrate. The QE increases within the range 77–175 K (probably due to reduced hole mobility), and next decreases in the higher temperature (probably due to influence of interfacial detects at the barrier/absorber interface). The insert in the figure for λ_{cut-off} = 3.4 μm shows that a low reverse
voltage is sufficient to reach maximum responsivity, where hB AlAsSb doping influence on the value of
nonzero turn-on voltage is visible.

![Figure 63](image1)

**Figure 63.** BSI spectral QE for MWIR detector for temperatures ranging from 77 to 225 K. The inset
shows the QE measured at 3.4 µm versus voltage at the same set of temperatures [171].

The MWIR T2SLs InAs/InAsSb nBn detector operating in HOT conditions has demonstrated
significant advantages over InSb. It is presented in Figure 64, where the 300 K background limited
detectivity versus detector temperature for f/2 optics and 2π field of view (FOV) is shown. The assessed
BLIP temperatures are 152 K and 181 K for f/2 and 2π FOV, respectively. In this way, this barrier detector
combines the higher operating temperature (comparable with HgCdTe) with the material robustness
of InSb. However, for low-background applications, the T2SLs InAs/InAsSb barrier detector still has
disadvantages compared to HgCdTe-higher G-R dark current due to shorter SRH lifetimes [171].

![Figure 64](image2)

**Figure 64.** 300 K background, 3-5 µm blackbody detectivity versus temperature for f/2 optics and
2π FOV (dashed lines); the corresponding ideal D* values are indicated by dotted lines. The shot noise
limited D* is also depicted (solid line) after [171]).

Hood et al. [192] modified the barrier structure to present better in performance pBn T2SLs
InAs/GaSb LWIR device. In the proposed architecture, the p-n junction can be located at the interface
between the heavily doped p-type contact and the low doped barrier or within the low doped barrier
itself. Similarly to the nBn, the pBn architecture reduces G-R currents related to the SRH centers
(the depletion region is moved to the barrier-no depletion in the low bandgap n-type absorber).
In addition, the electric field dropping in the barrier increases the response time which is related
to the carriers sweeping from the active layer. Recently, the design, growth, and characterization of the first MWIR T2SLs InAs/InAsSb pBn barrier photodetectors for HOT condition have been demonstrated [176,193].

In the last several years the idea of barrier detectors has been extended to short wavelength infrared wavelength (SWIR) region using InGaAs and InGaAsSb alloy systems [107,194–196]. The MBE technology is used as a standard growth method for SWIR detectors fabrication.

Savich et al. [195] carried out a comparison of electrical and optical performance of the typical photodiodes and nBn detectors with $\lambda_{\text{cut-off}} = 2.8 \, \mu m$ fabricated with both lattice-mismatched InGaAs and lattice-matched InGaAsSb active layers. In order to minimize the number of defects in the InGaAs active layer grown on InP substrate, a 2 $\mu m$ AlInAs step-graded buffer was grown where the lattice constant was graded from that of InP to that of In$_{0.82}$Ga$_{0.18}$As. Both the typical photodiode and nBn detector include this step-graded buffer. The extra pseudomorphic AlAsSb unipolar barrier is added to maintain a high barrier in conduction band in comparison to In$_{0.82}$Ga$_{0.18}$As.

In the case of lattice-matched to GaSb, the quaternary composition of In$_{0.30}$Ga$_{0.70}$As$_{0.56}$Sb$_{0.44}$ at the edge of the miscibility gap was used to maintain the $\lambda_{\text{cut-off}}$ and lattice-matching requirements. In nBn detector, also pseudomorphic AlGaSb unipolar barrier with a large CBO and zero VBO in comparison to the In$_{0.30}$Ga$_{0.70}$As$_{0.56}$Sb$_{0.44}$ absorber was implemented.

Figure 65 shows temperature dependent dark current characteristics for both lattice-mismatched InGaAs and lattice-matched InGaAsSb detectors under reverse bias 100 mV [195]. InGaAs on InP materials exhibit low material quality, i.e., threading dislocations occurring due to the lattice mismatch increasing dark current.

![Figure 65. Arrhenius plots for both InGaAs and InGaAsSb p-n junctions and nBn detectors with $\lambda_{\text{cut-off}} = 2.8 \, \mu m$ (adapted after [195]).](attachment:figure65.png)

The p-n InGaAs photodiode is limited by surface leakage current for $T < 220 \, \text{K}$, while the nBn detector remains diffusion limited down to 150 K. At the room temperature background photocurrent level, the nBn detector’s dark current is reduced by 400× in comparison to the typical photodiode.

The p-n InGaAsSb device is limited by depletion region current for $T < 250 \, \text{K}$, while the nBn remains diffusion limited down to 250 K. At the 300 K background photocurrent level of the nBn, dark current is suppressed by nearly three orders of magnitude in comparison to the typical photodiode.

Figure 66 shows that InGaAsSb nBn detector being lattice-matched to a GaSb substrate presents performance near “Rule 07” where dark current is 10 to 20 times lower in comparison to the lattice-mismatched InGaAs counterpart.
Figures 67 and 68 compare dark currents of different types of barrier and p-i-n detectors constructed on the basis of InAs/GaSb and Ga-free InAs/InAsSb SLs, respectively. Experimental data collected from the literature [107, 143, 147, 155, 161, 163, 173, 181, 196–202] is referred to the “Rule 07”. The dark currents of the InAs/GaSb SL detectors differ more from “Rule 07” than the data for Ga-free SL detectors. The best results were achieved by the pMp and the CBIRD detectors, both operating at 77 K. Fabricated by VIGO System Ga-free InAs/InAsSb SLs pBnN detector reaches dark currents measured at high temperatures (210 K and 230 K) in accordance with the “Rule 07” -see Figure 68. Similar results were reached by the detector described in the work of Kim et al. [163] but operating in 77 K. This figure also shows the advantage of barrier detectors over P-I-N devices.
6.2.3. Multicolor Detectors

Apart from HgCdTe photodiodes and quantum well infrared photodetectors (QWIPs), T2SLs have emerged as a candidate suitable for multispectral detection due to flexibility in bandgap tuning assuming lattice-matching requirements [203]. The single multicolor detector cell is built of the group of detectors being sensitive to a given spectral band (see Figure 69). The radiation goes through the SWIR detector, while the MWIR/LWIR passes freely to the next detector. The constituent layers absorb IR up to its $\lambda_{\text{cut-off}}$ being transparent to the longer wavelengths collected in subsequent layers. The detector structure is fabricated by proper order: MWIR/LWIR optically behind SWIR detector. The operating IR range is addressed by changing the polarity of the voltage.

![Figure 68. Comparison of various types Ga-free InAs/InAsSb SL detectors with the “Rule 07”.

Northwestern University presented different types of bias-switchable dual-band nBn T2SLs InAs/InAsSb detectors to include MWIR/LWIR and LWIR1/LWIR2 ranges [166,168,204]. The fabrication of dual-band photodetectors is challenging to grow proper electron barrier and SLs designs for both absorption regions with zero VB discontinuity. 

Figure 69 presents schematic diagram of a bias-selectable nBn dual-band IR detector based on T2SL InAs/InAsSb/AlAsSb structure together with band alignment of the SLs in two absorption layers. The SLs design of the red absorption region, with nominal $\lambda_{\text{cut-off}}$ ~12 µm at 77 K, consists of 30/10 MLs of T2SLs InAs/InAsSb0.48Sb0.52, respectively. The SLs design of the blue absorption region,
with a nominal λ\textsubscript{cut-off} ~8 µm at 77 K, consists of 13/1.5/13/9 MLs of T2SLs InAs/AlAs/InAs/InAs\textsubscript{0.48}Sb\textsubscript{0.52}, respectively. The SLs blue channel absorption region contains extra AlAs layer providing the necessary flexibility in the band structure design. The AlAs layer compensates the excessive tensile strain generated by the InAsSb layer in the SLs. The electron barrier is made of 4/3/4/3/4/9 MLs of T2SLs InAs/AlAs\textsubscript{0.35}Sb\textsubscript{0.45}/InAs/AlAs\textsubscript{0.45}Sb\textsubscript{0.55}/InAs/InAs\textsubscript{0.45}Sb\textsubscript{0.55} with a nominal λ\textsubscript{cut-off} ~4 µm at 77 K. This barrier design provides more flexibility to accommodate the lattice strain.

The two active regions are selected by changing the voltage polarity. The optical performance of the detector is shown in Figure 70a. The blue and red channels exhibit 100% λ\textsubscript{cut-off} ~8.6 µm (~145 meV) and ~12.5 µm (~100 meV) at 77 K, respectively. The blue channel QE saturates at the level of 65% at 6.45 µm and the red channel reaches 50% at 9 µm.

Figure 70. Performance characteristics of bias-selectable dual-band nBn InAs/InAsSb SL detector at 77 K: (a) spectral QE in FSI configuration without any anti-reflection coating for two applied bias: +40 mV (blue channel) and −100 mV (red channel), (b) dark current density versus voltage for blue and red channels, (c) D* of the blue and red channels at +40 and −100 mV voltage (after [168]).

Figure 70b shows the dark current density of the dual-band LWIR1/LWIR2 photodetector measured when covered by a 77K cold-shield. At +40 mV, dark current density for the blue channel reaches 1.13 × 10\textsuperscript{-6} A/cm\textsuperscript{2} and detectivity reaches 5.1 × 10\textsuperscript{12} cmHz\textsuperscript{1/2}/W at 77 K. The red channel exhibits dark current density at the level of 1.7 × 10\textsuperscript{-4} A/cm\textsuperscript{2} at −100 mV and detectivity of 4.5 × 10\textsuperscript{11} cmHz\textsuperscript{1/2}/W. The detectivity is estimated using Equation (15).

The results presented here indicate that the T2SLs InAs/InAsSb have the potential for multispectral IR imagers and can be a possible alternative for current state-of-the-art T2SLs InAs/GaSb.

7. InAsSb-based Superlattice Focal Plane Arrays

The noise equivalent difference temperature (NEDT) is a figure of merit for thermal imagers giving information on the detector array sensitivity and can be expressed according to the relations [3,78].

\[
\text{NEDT} = \frac{1 + \left( J_{\text{dark}} / J_{\Phi} \right)}{\sqrt{N_{w}} \left( \frac{1}{\Phi_{B}} \frac{d\Phi_{B}}{dT} \right)}
\]

\[
J_{\Phi} = \eta \Phi_{B} A
\]

\[
\sqrt{N_{w}} = \frac{\left( J_{\text{dark}} + J_{\Phi} \right) \tau_{\text{int}}}{q}
\]

where \( \Phi_{B} \)—background flux, \( \tau_{\text{int}} \)—integration time, \( N_{w} \)—well capacity of readout, \( J_{\Phi} = \eta \Phi_{B} A \)—background flux current. The overall QE of the detector, including the internal QE is given by \( \eta \). The cold shield efficiency and optics transmission are taken to be unity.
Assuming that the scene contrast, \( C = (d\Phi_B/dT)/\Phi_B \), \( \text{NEDT} \) can be rewritten:

\[
\text{NEDT} = \frac{1 + (I_{\text{dark}}/I_\Phi)}{\sqrt{N_C C}}. \tag{19}
\]

The calculated \( C \) for MWIR and LWIR is 3.5–4% and 1.7% at 300 K, respectively. Well capacity for a 15-\( \mu \)m pixel stays within the range \( N_w = 1 \times 10^6 \) to \( 1 \times 10^7 \) electrons. Equation (18) indicates that if the \( I_{\text{dark}}/I_\Phi \) ratio increases and/or \( \text{QE} \) decreases, the longer integration time and the faster optics have to be implemented.

Figure 71 presents \( \text{NEDT} \) versus temperature for barrier detectors and HgCdTe photodiodes with \( \lambda_{\text{cut-off}} = 5 \) \( \mu \)m and 10 \( \mu \)m. The theoretical HgCdTe photodiodes performance limit is higher than for barrier detectors for \( T > 150 \) K in a MWIR and \( T > 80 \) K in a LWIR range. However, taking into account both types of T2SLs, the operating temperature of InAs/InAsSb based detector is higher. For low temperatures the figure of merit of both T2SLs allows for similar performance to be reached, being mainly limited by the readout circuits.

7.1. InAsSb nBn Detector FPAs

The MWIR InAsSb nBn arrays are fabricated by several companies. The nBn cell is considered to be self-passivating design, decreasing leakage current and improving reliability and manufacturability. The simple design (see Table 5) is a major advantage in the large IR FPA state of the art.

First nBn InAsSb array commercially developed by Lockheed Martin Santa Barbara Focalplane operates within the temperature range \( T = 145–175 \) K. The 12 \( \mu \)m pixel-pitch MWIR nBn sensor of the 1280 \( \times \) 1024 format is packaged in a 1.4” diameter dewar with net housing length ~3.8” to include the cooler. That cooler and electronics are reported to consume 5 W and guarantees 25k hours of operation [205].

Klipstein et al. demonstrated nBn array operating in the MWIR (3.4–4.2 \( \mu \)m) at \( T = 150 \) K, with \( f/5.5 \) optics developed by SCD known as “Kinglet” [206]. The 640 \( \times \) 512 “Kinglet” is based on SCD’s Pelican-D readout integrated circuit (ROIC) and 15-\( \mu \)m pitch InAs\( _{0.91}\)Sb\( _{0.09}\)/B-AlAsSb architecture. The \( \text{NEDT} \) for \( f/3.2 \) optics and the pixel operability versus temperature is presented in Figure 72, reaching 20 mK at \( \tau_{\text{int}} = 10 \) ms and the pixel operability after a typical two point nonuniformity correction is higher than 99.5%, respectively. The \( \text{NEDT} \) and operability change sharply for \( T > 170 \) K, being consistent with the BLIP temperature of 175 K.
Recently, SCD has presented several 10 μm pitch nBn 1920 × 1536, 1280 × 1024 and 640 × 512 arrays, integrated with the Blackbird ROICs designed for HOT conditions [207]. Table 6 summarizes the performance of two InAsSb barrier devices fabricated by IR Cameras and SCD. In comparison with standard InSb photodiode arrays, the integrated detector cooler assembles (IDCAs) of nBn devices are characterized by SWaP advantages. For 15-μm pitch 1280 × 1024 FPA, the IDCA weight is reduced by 10%, the power consumption by 70%, and the mean time to failure (MTTF) is increased by more than 100%. Figure 73 presents an image reached by a HOT Hercules array at 150 K [208]. The very first demonstration of the 5 μm pixels 2040 × 1156 was reported in [209].

**Figure 72.** “Kinglet” NEDT and pixel operability temperature dependence for f/3.2 optics (after [206]).

| Parameter                  | Performance                        |
|----------------------------|------------------------------------|
| Array format               | 1280 × 1024                        |
| Pixel pitch                | 12 μm                              |
| FPA temperature            | 150 K                              |
| FPA spectral range         | 1–5.3 μm                           |
| NEDT                       | <35 mK                             |
| Well capacity              | 2 Me⁻                              |
| Maximum frame rate         | 119                                 |
| Optics                     | f/2.3, f/4                         |
| Cooler options             | 5 W                                |
| Weight                     | <7 pounds                          |
| Size                       | 5.1” × 5.8” × 8”                    |

**Table 6.** nBn InAsSb FPA characteristics.
Whereas the e-SWIR detector is based on a bulk InGaAsSb absorber, the T2SLs InAs/InAsSb active layers were used for MWIR and LWIR. Table 7 presents high-performance, cost-effective 640 × 512 FPAs for MWIR and LWIR ranges. HOT nBn photodetectors operate at 150 K covering the near infrared (NIR, 0.75–1.4 μm), SWIR (1.4–3 μm), and MWIR atmospheric transmission window (3–5 μm) with very high QE. Figure 74 shows 640 × 512 FPAs images taken in different spectral ranges.

### Table 7. Performance of InAs/InAsSb T2SLs nBn detector FPAs ([172,211]).

|                      | MWIR                      | LWIR                      |
|----------------------|---------------------------|---------------------------|
| **Format**           | 640 × 512                 | 640 × 512                 |
| **Pixel size**       | 24 × 24 μm²               | 24 × 24 μm²               |
| **Cut-off wavelength**| 5.37 μm (160 K)           | 12.5 μm (60 K)            |
| **Quantum efficiency**| 52% (150 K, no A/R coating) | 30% (60 K)               |
| **Dark current**     | 9.6 × 10⁻⁵ A/cm² (~0.2 V, 157 K) (~4.5 × Rule’07) | 2.6 × 10⁻⁵ A/cm² (~0.05 V, 60 K) |
| **Detectivity**      | 3 × 10¹¹ Jones (150 K, f/2 optics, 300 K background) | 16.3 mK (62 K)           |
| **NEDT**             | 18.7 mK (160 K); 26.6 mK (170 K) | 99.7% (60 K)             |

Figure 74. e-SWIR, MIWR, and LWIR images registered by Sb-based FPA. The e-SWIR FPA is based on a bulk InGaAsSb, T2SLs InAs/InAsSb active layer was used for the MWIR and LWIR (after [107]).

Figure 75 shows the performance of the 24 μm pitch 640 × 512 FPA with $\lambda_{cut-off} \sim 5.4$ μm (see Table 7) and direct injection (DI) ROIC with $N_{e0} = 8 \times 10^9$ electrons. NEDT is below 19 mK for $T < 160$ K, and increases up to $\sim 55$ mK at 180 K. The operability is 99.7% up to 160 K and decreases to 99.3% at 180 K. NEDT exhibits excellent uniformity with 8 mK distribution [172]. Both NEDT and operability...
are similar to those described previously for 15 μm pitch nBn InAs$_{0.9}$Sb$_{0.09}$/B-AlAsSb 640 × 512 FPA shown in Figure 72.

The devices were grown on a 4", Te-doped GaSb (100) substrate by MBE. The LWIR barrier structures have also shown significant improvement in operating temperature in comparison to the typical p-n designs. Figure 76 presents the dark current density and the QE versus temperature for LWIR square mesa T2SLs InAs/InAsSb barrier detectors (area 250×250 μm$^2$) [212]. The devices were grown on a 4", Te-doped GaSb (100) substrate by MBE. The −50 mV dark current density at 70 K is $4 \times 10^{-9}$ A/cm$^2$. The QE derived from BSI (through the GaSb substrate) responsivity measurements has not been corrected for substrate reflection or transmission.

Figure 76. NEDT and operability versus temperature for 24 μm pitch MWIR 640 × 512 nBn InAs/InAsSb FPA.

The LWIR detector was used to fabricate 24 μm pitch, 640 × 512 FPA and Lockheed Martin SBF-193 ROIC. The measured 640 × 512 FPA NEDT histogram for $T = 60$ K, 300 K blackbody temperature and f/4 cold stop is shown in Figure 76c. The 21 mK NEDT is in fair agreement with the estimated NEDT based on a single element test detector data [212].

8. Conclusions

There are many critical challenges for future civilian and military IR detector applications. The main efforts are directed to decrease IR imaging systems SWaP by increasing the FPA’s operating temperature. Nowadays, HgCdTe is the most common material for IR photodetectors being the reference for alternative technologies to include uncooled operation. After sixty years of development, HgCdTe has just reached ultimate HOT performance limit estimated by “Law 19”-the ultimate photodiode performance metric [117]. This metric has been recently supported by experimental data
of fully-depleted HgCdTe P-i-N photodiodes. The “Law 19” corresponds exactly the BLIP for room temperature operation.

The InAsSb-based IR photodetectors’ performance is comparable with HgCdTe:

- III-V compounds exhibit inherently short SRH lifetimes typically below 1 µs requiring the nBn design to operate at higher temperatures (this applies both to the bulk InAsSb and the T2SLs),
- T2SLs InAs/InAsSb has emerged in recent years as an alternative to the T2SLs InAs/GaSb due to the three reasons: (i) a better growth controllability and simpler manufacturability, (ii) longer SRH lifetimes (10 µs in MWIR SLs in comparison with 400 ns for InSb), and (iii) significantly higher operating temperature than InSb,
- HgCdTe exhibits long SRH lifetimes >200 µs to 50 ms depending on the \( \lambda_{\text{cut-off}} \) \[78\] (can operate with either architecture and may be diffusion or depletion current limited),
- III-Vs exhibit comparable performance to HgCdTe for equivalent \( \lambda_{\text{cut-off}} \), but for lower temperature, due to the inherent difference in SRH lifetimes,
- III-V materials are more robust than their II-VI counterparts due to stronger (less ionic) chemical bonding which allows to reach better operability, spatial uniformity, temporal stability, scalability, producibility, and affordability-the so-called “ibility” advantages.

Table 8 gathers the LWIR detectors state of the art fabricated from selected materials. The highest level of maturity (technology readiness level, TRL = 9) is credited to HgCdTe photodiodes, microbolometers, and for QWIPs. The T2SLs InAs/InAsSb structures have proven the potential for MWIR and LWIR applications with performance comparable to HgCdTe for the same \( \lambda_{\text{cut-off}} \).

Strong progress toward mature SLs and barrier detector technologies, including their commercialization, has been observed in the last decade. It is expected that InAsSb-based materials strengthen dominant position of InSb especially in MWIR FPA market in volume due to “ibility” advantages of III-V semiconductors. HgCdTe detectors allow one to reach lower dark current and HOT conditions and are/will be a material of choice for more demanding applications.

Table 8. LWIR device state-of-the-art. TRL-technology readiness level (TRL = 10 ideal maturity).

| Maturity level | Bolometer | HgCdTe | QWIP | T2 SLs InAsGaSb | InAsSb-Based |
|----------------|-----------|--------|------|-----------------|-------------|
| Status          | Applications where medium to low performance is required. | Application where high performance is required. | Commercial | Research and development | Research and development |
| Operating temp. | Un-cooled | Cooled | Cooled | Cooled | Cooled/TE cooled |
| Manufacturability | Excellent | Poor | Excellent | Very good | Very good |
| Cost | Low | High | Medium | Medium | Medium |
| Prospect for large format | Excellent | Very good | Excellent | Excellent | Excellent |
| Availability of large substrate | Excellent | Poor | Excellent | Very good | Very good |
| Military applications | Weapon sight, night vision goggles, missile seekers, small UAV sensors, unattended ground sensors. | Missile intercept, tactical ground and air born imaging, hyper spectral, missile seeker, missile tracking, space based sensing. | Being evaluated for some military applications and astronomy sensing. | Being evaluated for some military applications. | Being developed in university and evaluated industry research environment. |
| Limitations | Low sensitivity and long time constants, mechanical stability. | Performance susceptible to manufacturing variations. Difficult to extend to >14-\( \mu \)m cutoff. | Narrow bandwith and low sensitivity. | Requires a further investment and fundamental material breakthrough to mature. | Degradation of detector performance in a space environment due to the radiation-induced minority carrier lifetime degradation. |
| Advantages | Low cost and requires no active cooling, leverages standard Si manufacturing equipment. | Near theoretical performance, will remain material of choice for minimum of the next several years. High performance of substrate-removed arrays for space missions. | Low cost applications. Leverages commercial manufacturing processes. Very uniform material. | Theoretically better then HgCdTe, leverages commercial III-V fabrication techniques. | MWIR devices with significantly higher operating temperature than InSb. |
Author Contributions: Conceptualization, A.R., P.M. (Piotr Martyniuk); writing—original draft preparation, A.R., P.M. (Piotr Martyniuk), M.K., P.M. (Pawel Madejczyk), S.K.; writing—review and editing, A.R., P.M. (Piotr Martyniuk), M.K.; visualization, A.R., P.M. (Piotr Martyniuk), M.K.; supervision, A.R., P.M. (Piotr Martyniuk); project administration, A.R., P.M. (Piotr Martyniuk), M.K., P.M. (Pawel Madejczyk); funding acquisition, A.R., P.M. (Piotr Martyniuk), M.K., P.M. (Pawel Madejczyk). All authors have read and agreed to the published version of the manuscript.

Funding: The writing of the paper has been partially done under financial support of The National Science Centre-Grant nos. UMO-2018/30/M/ST7/00174, UMO-2018/31/B/ST7/01541, and UMO-2019/33/B/ST7/00614 and The National Centre for Research and Development—the grant no. Mazowsze/0090/19-00.

Conflicts of Interest: The authors declare no conflict of interest. The funders had no role in the design of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript, or in the decision to publish the results.

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