

InGaAs 自适应阱簇复合结构的偏振双峰辐射机制及能带特征

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摘要 研究一种基于富镉团簇(IRC)效应的新型高应变 InGaAs/GaAs 自适应阱簇复合(WCC)量子结构,该结构具有比较灵活的能带调控能力,可产生一种特殊的偏振双峰光谱。为探究该 WCC 结构的偏振双峰辐射机制及能带特征,利用 IRC 效应生长获得了自适应 WCC 结构,并测量了该新型结构的偏振光致发光(PL)光谱,在横向电场(TE)和横向磁场(TM)模式下,PL 光谱皆显示出特殊的双峰结构。这主要是镉原子的自适应迁移导致有源层同时存在镉组分正常和镉组分降低的 $\text{In}_x\text{Ga}_{1-x}\text{As}$ 材料所致。根据半导体激光器的跃迁矩阵元理论和 PL 光谱的双峰能量间距,阐明不同偏振模式下的光谱双峰与多组分 $\text{In}_x\text{Ga}_{1-x}\text{As}$ 材料的对应辐射机制,同时确定该 WCC 结构的混合能带特征。该研究内容对 IRC 效应和新型 WCC 发光结构的发展和应用具有重要研究意义。

关键词 半导体激光器; InGaAs/GaAs; 富镉团簇; 偏振双峰光谱; 混合能带

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1 引言

常规的 InGaAs/GaAs 应变量子阱具有固定的镉含量和单一的应变类型^[1-4],已经广泛应用于半导体光放大器、光纤通信、生物医学等众多领域^[5-7]。在生长常规的高应变 InGaAs 量子阱时,富镉团簇(IRC)效应通常被作为一种生长缺陷来规避^[8-10],因此忽略了其特殊的光学特性。所谓 IRC 效应,即在生长高应变 InGaAs/GaAs 量子结构时,为了释放 InGaAs 材料中积累的高应变,镉原子会沿着材料生长方向自动向上迁移,并在 InGaAs 材料表面堆积,形成三维团簇(IRC)。近几年,本课题组基于 IRC 效应,生长获得了一种新型非对称 InGaAs 自适应阱簇复合(WCC)量子限制结构,该 WCC 有源层包含了大量不同镉含量的 $\text{In}_x\text{Ga}_{1-x}\text{As}$ 激活区以及混合应变类型,且偏振光致发光(PL)谱具有明显的双峰结构。本课题组通过测量 WCC 样品两端的 PL 光谱,计算并分析了其超宽的增益特性,有望实现 100 nm 的超宽调谐激光器^[11]。此外,利用高斯拟合探究了 PL 光谱的构成,分析并实现了在单一 WCC 结构上的双波长的同步激励^[12]。同时,该 WCC 结构中几乎等大

的横向电场(TE)和横向磁场(TM)偏振增益对实现偏振独立光放大器也具有重要的研究价值^[13]。由于该 WCC 结构是基于 IRC 效应形成的,该效应会引起镉原子的自适应迁移,进而导致在单一 InGaAs 有源层内同时产生镉组分正常的和镉组分降低的 $\text{In}_x\text{Ga}_{1-x}\text{As}$ 区域以及张压应变同时存在的混合能带类型,以上复杂特征导致偏振双峰与多组分 $\text{In}_x\text{Ga}_{1-x}\text{As}$ 材料的对应辐射机制尚不清晰。因此,探究清楚 InGaAs 自适应 WCC 结构的偏振双峰辐射机制和混合能带特征对研究和评估新型 WCC 半导体激光器具有重要意义^[14-15]。

本文首先利用 IRC 效应生长获得了 InGaAs 自适应 WCC 量子结构,并在室温下测量了该结构在 TE 和 TM 模式下的 PL 光谱,其皆呈现出明显的两个谱峰。根据半导体激光器的跃迁矩阵元理论,阐明了偏振光谱双峰与多组分 $\text{In}_x\text{Ga}_{1-x}\text{As}$ 材料的对应辐射机制,同时获得了包含第一导带(C_1)以及第一重空穴带(HH_1)和轻空穴带(LH_1)的混合能带结构。

2 $\text{In}_x\text{Ga}_{1-x}\text{As}$ 自适应 WCC 量子结构

利用金属有机化学气相沉积系统(MOCVD),在

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未掺杂的 GaAs(001) 衬底上生长获得了基于 IRC 效应的 InGaAs 自适应 WCC 量子结构。形成 IRC 的基本原理是在生长高应变 $\text{In}_x\text{Ga}_{1-x}\text{As}/\text{GaAs}$ 量子结构时, 为了释放 $\text{In}_x\text{Ga}_{1-x}\text{As}$ 量子阱中逐渐积累的高应变, 镉原子会沿着材料生长方向自动向上迁移, 并在 $\text{In}_x\text{Ga}_{1-x}\text{As}$ 材料表面堆积, 形成富镉团簇^[16-17]。为了生长获得 InGaAs 自适应 WCC 量子结构, 本文设计 $\text{In}_{0.17}\text{Ga}_{0.83}\text{As}/\text{GaAs}/\text{GaAsP}_{0.08}$ 材料体系为有源区, 其中 $\text{GaAsP}_{0.08}$ 势垒层厚 8 nm, 用来吸收泵浦光; $\text{In}_{0.17}\text{Ga}_{0.83}\text{As}$ 势阱的层厚被设计为 10 nm; 由于 WCC 量子结构的形成主要取决于 $\text{In}_{0.17}\text{Ga}_{0.83}\text{As}$ 材料内部的应变积累, 为了更好地标定和补偿 $\text{In}_{0.17}\text{Ga}_{0.83}\text{As}$ 材料内部的应变, 在势阱和势垒之间增加了 2 nm 厚的 GaAs

应变补偿(SC)层。InGaAs 自适应 WCC 量子限制结构如图 1(a) 所示。其中 10 nm 的厚度设计是为了获得足够的应变积累, 有利于 IRC 的产生。只有当 $\text{In}_x\text{Ga}_{1-x}\text{As}$ 层的生长厚度超过若干个单分子层后, 为释放较大晶格失配引起的应变积累, 镉原子才会迁移到 InGaAs 层表面堆积并形成 IRC, 同时会在有源区产生镉组分正常的 $\text{In}_{0.17}\text{Ga}_{0.83}\text{As}$ 材料和镉组分降低的 $\text{In}_x\text{Ga}_{1-x}\text{As}$ 区域, 形成原理如图 1(b) 所示。为确保生长的 WCC 结构中包含 IRC, 本文使用原子力显微镜 (AFM; XE100, Park Systems Instrument 公司) 测量获得了 InGaAs 的上表面形貌, 结果如图 1(a) 中的插图所示。原子力照片中展现了大量不规则的三维团簇形貌, 即 IRC, 说明该结构产生了 IRC 效应。

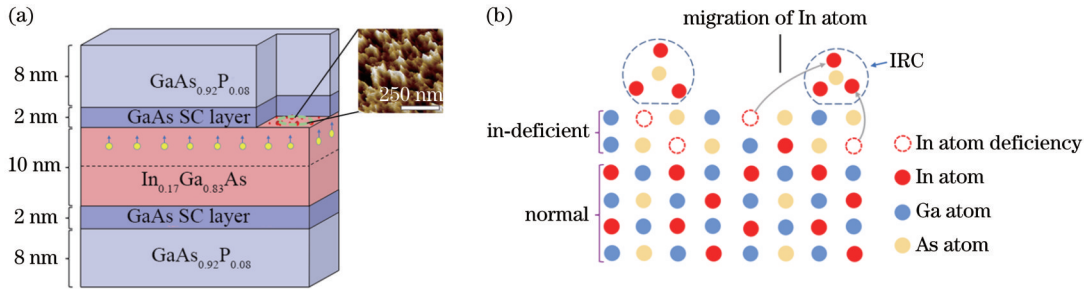


图 1 InGaAs 自适应 WCC 量子结构及 IRC 形成机制。(a) WCC 量子结构及 IRC 形貌; (b) IRC 形成原理

Fig. 1 InGaAs self-fit WCC quantum structure and formation mechanism of IRC. (a) WCC quantum structure and IRC topography; (b) formation principle of IRC

3 偏振双峰光谱的实验测量

为了探究 InGaAs/GaAs WCC 量子结构的偏振双峰辐射机制, 在不同的载流子注入浓度下 ($N=3.0 \times 10^{17}, 3.6 \times 10^{17}, 3.9 \times 10^{17}, 4.5 \times 10^{17} \text{ cm}^{-3}$), 利用线性偏振片分别测量了该 WCC 结构的 TE 和 TM 偏振 PL

光谱, 测量原理如图 2(a) 所示。为避免端面反射对光谱峰值的影响, 该 WCC 结构两端都进行了增透处理, 透过率 $T=99.99\%$ 。波长为 808 nm 的光纤耦合激光器作为泵浦源, 线性偏振片和两个焦距为 10 cm 的平凸透镜用于收集不同偏振模式下的 PL 谱, 测量结果如图 2(b) 所示。

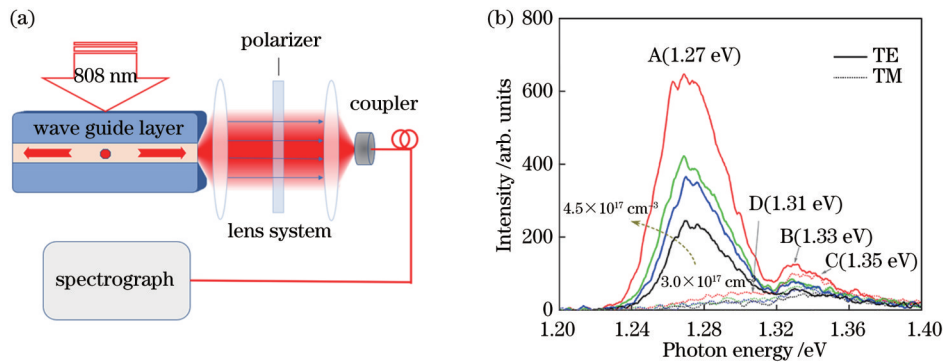


图 2 PL 光谱的测量原理及光谱曲线。(a) PL 光谱的测量原理; (b) TE 和 TM 偏振模式下的 PL 光谱

Fig. 2 PL spectral measurement principle and spectral curves. (a) PL spectral measurement principle; (b) PL spectra in the TE and TM polarizations

图 2(b) 中 TE 和 TM 偏振 PL 光谱皆呈现出特殊的双峰特征, 即每条谱线都具有明显的两个峰, 这里分别用 A、B 和 C、D 标记两种偏振模式下的峰值点, 对应的光子能量分别为 1.27 eV、1.33 eV 和 1.35 eV、

1.31 eV, 峰谷能量约为 1.317 eV。该双峰特征产生的原因是在 $\text{In}_x\text{Ga}_{1-x}\text{As}/\text{GaAs}$ 材料生长过程中发生了 IRC 效应, 镉原子的自适应迁移导致正常 $\text{In}_{0.17}\text{Ga}_{0.83}\text{As}$ 材料中镉含量降低, 形成镉组分降低的 $\text{In}_x\text{Ga}_{1-x}\text{As}$ 区

域。由于该低钢区域带隙较宽, PL 谱峰向短波方向偏移。该低钢区域与正常的 $\text{In}_{0.17}\text{Ga}_{0.83}\text{As}$ 材料共同产生了图 2 中的双峰光谱。由于该 WCC 结构是基于 IRC 效应产生的, 钢原子因自适应迁移会在顶部堆积, 产生 IRC 以及钢组分降低的有源区。富钢团簇的大小和分布具有一定的随机性, 会导致有源区的钢含量产生一定起伏, 进而会影响光谱的细节特征, 但总体的双峰特征不变^[11-12]。为探究 TE 和 TM 偏振光谱双峰与多组分 $\text{In}_x\text{Ga}_{1-x}\text{As}$ 区域的对应辐射机制及能带特征, 本文将利用跃迁矩阵元理论和 Model-Solid 理论进行分析和验证。

4 分析与讨论

4.1 WCC 结构的能带特征

根据 Model-Solid 理论, 当载流子注入浓度较低时, 电子主要占据带边能级, 光谱峰值对应的光子能量主要取决于导带 C_1 到价带之间的能量间隔。根据本课题组的前期研究, 该非对称 WCC 结构主要包括两种不同钢含量的 $\text{In}_x\text{Ga}_{1-x}\text{As}$ 材料, 即钢组分降低的

$\text{In}_{0.12}\text{Ga}_{0.88}\text{As}$ 层和钢组分正常的 $\text{In}_{0.17}\text{Ga}_{0.83}\text{As}$ 材料^[13]。由于钢原子的自适应迁移是在 $\text{In}_{0.17}\text{Ga}_{0.83}\text{As}$ 生长到若干个单分子层后发生的, 因此, 低钢的 $\text{In}_{0.12}\text{Ga}_{0.88}\text{As}$ 材料位于正常的 $\text{In}_{0.17}\text{Ga}_{0.83}\text{As}$ 层上表面, 如图 1 所示。为确定该非对称量子结构的应变类型, $\text{In}_x\text{Ga}_{1-x}\text{As}$ 材料的晶格常数 $a(x)$ ^[18] 可表示为

$$a(x) = 6.0583 - 0.405(1 - x), \quad (1)$$

式中 x 表示 $\text{In}_x\text{Ga}_{1-x}\text{As}$ 材料的钢含量。根据式 (1) 求得 GaAs、 $\text{In}_{0.17}\text{Ga}_{0.83}\text{As}$ 和 $\text{In}_{0.12}\text{Ga}_{0.88}\text{As}$ 的晶格常数分别为 5.65325 \AA 、 5.72215 \AA 和 5.7019 \AA ($1 \text{ \AA} = 10^{-10} \text{ m}$), 其晶格失配情况如图 3(a) 所示。经计算可知, 生长在 GaAs 上的 $\text{In}_{0.17}\text{Ga}_{0.83}\text{As}$ 材料和生长在 $\text{In}_{0.17}\text{Ga}_{0.83}\text{As}$ 上的 $\text{In}_{0.12}\text{Ga}_{0.88}\text{As}$ 材料对应的应变类型分别为 0.012 和 -0.004 。因此, 这种特殊的自适应 WCC 量子限制结构产生了一种特殊的混合应变类型, 即在正常的 $\text{In}_{0.17}\text{Ga}_{0.83}\text{As}$ 层中产生压应变, 其重空穴带 HH_1 位于轻空穴带 LH_1 上方; 在低钢的 $\text{In}_{0.12}\text{Ga}_{0.88}\text{As}$ 层中产生张应变, 其轻空穴带 LH_1 位于重空穴带 HH_1 上方。混合能带结构如图 3(b) 所示。

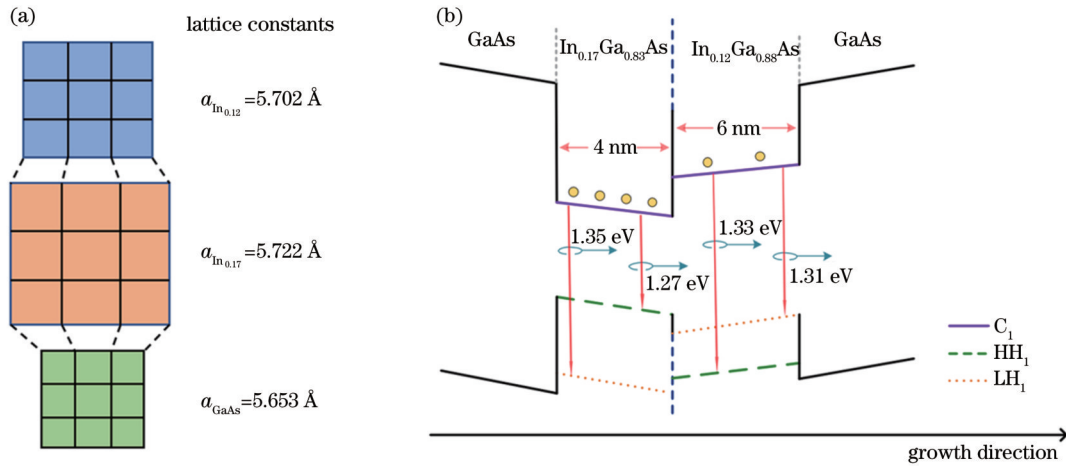


图 3 $\text{In}_x\text{Ga}_{1-x}\text{As}/\text{GaAs}$ 材料系统的晶格失配及 InGaAs 自适应 WCC 量子结构的能带特征。(a) $\text{In}_{0.12}\text{Ga}_{0.88}\text{As}$ 、 $\text{In}_{0.17}\text{Ga}_{0.83}\text{As}$ 与 GaAs 材料的晶格失配; (b) 能带特性

Fig. 3 Lattice mismatch of $\text{In}_x\text{Ga}_{1-x}\text{As}/\text{GaAs}$ material system and energy band characteristics of InGaAs self-fit WCC quantum structure. (a) Lattice mismatch of $\text{In}_{0.12}\text{Ga}_{0.88}\text{As}$, $\text{In}_{0.17}\text{Ga}_{0.83}\text{As}$, and GaAs; (b) energy band characteristic

4.2 偏振双峰的对应辐射机制

为探究偏振光谱的双峰与混合能带的对应辐射机制, 根据半导体激光器的跃迁矩阵元理论进行如下分析。当电子在导带 C_1 和重空穴 HH_1 能级之间跃迁时, TE 偏振的跃迁矩阵元值为 $|M|_{\text{TE}}^2 = |M|^2/2$, 而 TM 偏振的跃迁矩阵元 $|M|_{\text{TM}}^2 = 0$, 此时产生 TE 偏振的电子跃迁速率大于 TM 偏振的电子跃迁速率, TE 偏振的光谱强度会明显大于 TM 偏振光谱; 而当电子从导带 C_1 跃迁到轻空穴 LH_1 时, TE 偏振跃迁矩阵元 $|M|_{\text{TE}}^2 = |M|^2/6$, 而 TM 偏振跃迁矩阵元 $|M|_{\text{TM}}^2 = 2|M|^2/3$, TM 偏振的光谱强度呈现出超过 TE 偏振光谱的趋势,

其中的 $|M|^2$ 是一个常量, 为材料的动量矩阵元^[19]。因此, TE 偏振光谱主要由 C_1 和重空穴 HH_1 之间的电子-空穴复合决定, TM 偏振光谱主要由 C_1 与轻空穴 LH_1 之间的电子-空穴复合决定, TE 和 TM 光谱峰值之间的能量间隔对应于 HH_1 和 LH_1 之间的能量间隔。

根据跃迁矩阵元理论, 在压应变的 $\text{In}_{0.17}\text{Ga}_{0.83}\text{As}$ 材料中, 其重空穴带 HH_1 位于轻空穴带 LH_1 上方, 该材料辐射的 TE 偏振的光子能量要小于 TM 偏振光谱。而在具有张应变的 $\text{In}_{0.12}\text{Ga}_{0.88}\text{As}$ 材料中, 其轻空穴带 LH_1 位于重空穴带 HH_1 上方, TM 光谱对应的光子能量要小于 TE 光谱。根据跃迁矩阵元的分析, 并结合图 2 (b) 中的不同偏振模式下的双峰可以判断: TE 和 TM

光谱曲线中的主峰 A(1.27 eV)和 C(1.35 eV)主要是压应变 $\text{In}_{0.17}\text{Ga}_{0.83}\text{As}$ 材料发生辐射产生的;图 2(b)中 TE 和 TM 曲线中的次峰 B(1.33 eV)和 D(1.31 eV)主要是张应变的 $\text{In}_{0.12}\text{Ga}_{0.88}\text{As}$ 材料发生辐射产生的。偏振双峰的对辐射机制及混合能带特征如图 3(b)所示。另外,图中正常的 $\text{In}_{0.17}\text{Ga}_{0.83}\text{As}$ 材料厚度为 4 nm,这一结论是基于本课题组前期对 WCC 结构进行逐点扫描测量得到的实验结果^[20]。

通过图 3(b)可知:对于压应变 $\text{In}_{0.17}\text{Ga}_{0.83}\text{As}$ 材料,其重空穴和轻空穴的能量间距为 0.08 eV;对于张应变 $\text{In}_{0.12}\text{Ga}_{0.88}\text{As}$ 材料,重轻空穴的能量间隔为 0.02 eV。为验证 WCC 结构的能带特性及偏振双峰对应辐射机制的准确性,根据 Model-Solid 理论分别计算得到了压应变 $\text{In}_{0.17}\text{Ga}_{0.83}\text{As}$ 材料和张应变 $\text{In}_{0.12}\text{Ga}_{0.88}\text{As}$ 材料中 LH_1 和 HH_1 之间的能量间隔^[21-22],计算公式为

$$\Delta E_{\text{HH}}(x, y) = 2a_v \left(1 - \frac{C_{12}}{C_{11}}\right) \epsilon + |\mathbf{b}| \left(1 + 2 \frac{C_{12}}{C_{11}}\right) \epsilon, \quad (2)$$

$$\Delta E_{\text{LH}}(x, y) = 2a_v \left(1 - \frac{C_{12}}{C_{11}}\right) \epsilon - |\mathbf{b}| \left(1 + 2 \frac{C_{12}}{C_{11}}\right) \epsilon, \quad (3)$$

式中: ΔE_{HH} 和 ΔE_{LH} 分别代表应变引起的重空穴和轻空穴的能带偏移量; a_v 表示半导体材料中价带的形变势; ϵ 表示应变; C_{11} 和 C_{12} 代表弹性常量; \mathbf{b} 代表柏氏向量。根据式(2)和式(3)可知,晶格失配引起的应变会导致 HH_1 和 LH_1 发生分离和偏移。在 $\text{In}_{0.17}\text{Ga}_{0.83}\text{As}$ 材料中, HH_1 和 LH_1 的能带偏移分别为 $\Delta E_{\text{HH}} = 0.025$ eV 和 $\Delta E_{\text{LH}} = -0.0544$ eV,因此 HH_1 和 LH_1 之间的能量间隔为 $\Delta E_{0.17} = 0.079$ eV。对于 $\text{In}_{0.12}\text{Ga}_{0.88}\text{As}$ 材料, HH_1 和 LH_1 的能带偏移分别为 $\Delta E_{\text{HH}} = -0.0073$ eV 和 $\Delta E_{\text{LH}} = 0.0161$ eV,因此 HH_1 和 LH_1 之间的能量间隔为 $\Delta E_{0.12} = 0.023$ eV。理论结果与图 3 所示的实验数据基本一致。这表明了该非对称 InGaAs 自适应 WCC 量子结构中能带特征及偏振双峰对应辐射机制的准确性。

5 结 论

基于生长高应变 InGaAs 量子阱中的富镉团簇效应,生长获得了一种特殊的非对称自适应 WCC 结构,测量了该新型结构 TE 和 TM 模式下的光致发光光谱,光谱皆呈现出特殊的双峰特征。根据半导体激光器的跃迁矩阵元理论,结合偏振双峰的能量间距,阐明了偏振双峰与多组分 $\text{In}_x\text{Ga}_{1-x}\text{As}$ 材料的对应辐射机制,同时确定了该非对称结构的能带具有压/张应变同时存在的混合应变特征。该研究结果对分析 InGaAs 非对称 WCC 结构具有重要意义。

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Polarized Dual-Peak Radiation Mechanism and Energy-Band Characteristics of InGaAs Self-Fit Well-Cluster Composite Structure

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Abstract

Objective In this paper, a novel, highly-strained InGaAs/GaAs self-fit well-cluster composite (WCC) quantum structure was investigated. The WCC structure differs from the conventional InGaAs/GaAs quantum-well structure, which exhibits considerable potential for numerous laser applications. Presently, the conventional quantum well exhibits a quasirectangle well structure, wherein each well consists of a fixed amount of indium and a single strain type in the material system. The WCC structure with variable indium content and thickness in an $\text{In}_x\text{Ga}_{1-x}\text{As}/\text{GaAs}$ system can yield remarkable results, thereby facilitating the development of new laser types. This structure is associated with the indium-rich cluster (IRC) effect, wherein the IRCs were typically regarded as defects to be avoided for the conventional InGaAs quantum-well structure; hence, its special optical characteristics remain neglected. The migration of the indium atoms to the WCC structure would reduce the indium content in the corresponding InGaAs regions, consequently generating normal and indium-deficient $\text{In}_x\text{Ga}_{1-x}\text{As}$ regions with hybrid strain types in the InGaAs material and aid in the production of special polarized spectra with dual peaks. Therefore, it is crucial to reveal the underlying corresponding luminescence mechanism between dual peaks in different polarized spectra and multiple $\text{In}_x\text{Ga}_{1-x}\text{As}$ materials. This work offers new avenues for the development of new types of devices.

Methods First, the InGaAs-based WCC quantum structure was developed via metal-organic chemical vapor-phase deposition. To generate the IRC effect by sufficient strain accumulation, the $\text{In}_{0.17}\text{Ga}_{0.83}\text{As}/\text{GaAs}/\text{GaAsP}_{0.08}$ material system was designed as a periodic gain structure (Fig. 1). The thickness of the $\text{In}_{0.17}\text{Ga}_{0.83}\text{As}$ layer was designed to be 10 nm, because an InGaAs layer thinner than 10 nm is insufficient to obtain the IRC effect. Second, the luminescence mechanism of the WCC structure was studied by coating the WCC sample at a transmittance of $T=99.99\%$ at the dual facets to avoid the end reflection. Third, the polarized photoluminescence (PL) spectra in transverse electric (TE) and transverse magnetic (TM) modes were measured using a linear polarizer under varying carrier densities (N) in the range of $3.6 \times 10^{17} - 4.8 \times 10^{17} \text{ cm}^{-3}$. The special bimodal feature was observed in both TE- and TM-polarized PL spectra (Fig. 2), which could be associated with the emissions from normal and indium-deficient $\text{In}_{0.12}\text{Ga}_{0.88}\text{As}$ regions with varying band gaps. To analyze the different strain types, the lattice constant $a(x)$ of the $\text{In}_x\text{Ga}_{1-x}\text{As}$ material was obtained. Subsequently, the hybrid strain types in normal and indium-deficient $\text{In}_x\text{Ga}_{1-x}\text{As}$ were obtained. The compressive and tensile strain occurred in the $\text{In}_{0.17}\text{Ga}_{0.83}\text{As}$ and $\text{In}_{0.12}\text{Ga}_{0.88}\text{As}$ layers, respectively. Further, according to the transition matrix element theory, the PL spectrum in TE polarization was primarily associated with the electron-hole recombination

between the first conduction (C_1) and heavy hole (HH_1) subbands, whereas that in TM polarization was associated with the electron-hole recombination between C_1 and light hole (LH_1) subbands. Moreover, in the compressively-strained $In_{0.17}Ga_{0.83}As$ layer, the HH_1 subband lies above the LH_1 subband. Conversely, the HH_1 subband lies below the LH_1 subband in the tensile-strained $In_{0.12}Ga_{0.88}As$ layer. Finally, the underlying luminescence mechanism of the polarized spectra with dual peaks was revealed according to the above analysis.

Results and Discussions The TE- and TM-polarized PL spectra (Fig. 2) reveal the special bimodal features, and they are marked using letters A and B, and C and D, which correspond to 1.27 eV and 1.33 eV and 1.35 eV and 1.31 eV, respectively. GaAs, $In_{0.17}Ga_{0.83}As$, and $In_{0.12}Ga_{0.88}As$ yield lattice constant values of 5.65325, 5.72215, and 5.7019 Å, respectively. Accordingly, the $In_{0.17}Ga_{0.83}As$ layer is subject to compressive strain such that the HH_1 subband lies above the LH_1 subband. Meanwhile, tensile strain is noted in the $In_{0.12}Ga_{0.88}As$ layer such that the HH_1 subband falls below the LH_1 subband. According to the transition matrix element theory, the main peaks, which are marked by A and C in both TE and TM spectral curves (Fig. 2), are attributed to the compressively-strained $In_{0.17}Ga_{0.83}As$ layer; the corresponding TE photon energy is less than the TM photon energy. Meanwhile, the subpeaks marked by B and D (Fig. 2) are attributed to the tensile-strained $In_{0.12}Ga_{0.88}As$ region; the corresponding TE photon energy exceeds the TM photon energy. Moreover, the characteristics of the hybrid energy band of the InGaAs self-fit WCC quantum structure are obtained (Fig. 3). For the compressively-strained $In_{0.17}Ga_{0.83}As$ layer, the energy intervals from C_1 to HH_1 and LH_1 bands are 1.27 eV and 1.35 eV, which correspond to the photon energy at peaks A and C, respectively. In the case of the tensile-strained $In_{0.12}Ga_{0.88}As$ material, the energy intervals from C_1 to HH_1 and LH_1 bands are 1.33 eV and 1.31 eV, which correspond to the photon energy at peaks B and D, respectively.

Conclusions In this paper, a novel, highly-strained InGaAs/GaAs self-fit WCC quantum structure is investigated based on the IRC effect. The measured spectra in TE and TM polarizations reveal special features with dual peaks, which are attributed to the combination of different emissions produced by the normal and indium-deficient InGaAs active regions. According to the transition matrix element theory, the corresponding luminescence mechanism between the dual peaks in polarized spectra and InGaAs material with different indium content is revealed. Furthermore, the band characteristics associated with the conduction subband C_1 and valence subbands of heavy holes HH_1 and light holes LH_1 are determined to reveal the underlying luminescence mechanism. The special WCC structure demonstrates a hybrid strain distribution, which indicates the simultaneous existence of compressive and tensile strains. The results of this study can greatly enhance the performance of InGaAs-based WCC-tunable lasers.

Key words semiconductor laser; InGaAs/GaAs; indium-rich cluster; polarized dual-peak spectrum; hybrid energy band