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(54) METHODS OF FORMING SINGLE SOURCE PRECURSORS, METHODS OF FORMING POLYMERIC SINGLE SOURCE PRECURSORS, AND SINGLE SOURCE PRECURSORS AND INTERMEDIATE PRODUCTS FORMED BY SUCH METHODS

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C07F 1/00 (2006.01)

- (52) **U.S. Cl.** **556/14**; 556/19; 556/110

(56) References Cited

U.S. PATENT DOCUMENTS

4,687,881	A	8/1987	Goslowsky et al.
4,906,290	A	3/1990	Worner
5,445,847	Α	8/1995	Wada et al.
5,501,786	A	3/1996	Gremion et al.
5,567,469	A	10/1996	Wada et al.
5,858,120	A	1/1999	Nakagawa et al.
6,127,202	Α	10/2000	Kapur et al.
6,145,342	Α	11/2000	Bayya et al.
6,307,148	В1	10/2001	Takeuchi et al.
6,355,874	В1	3/2002	Yagi et al.
6,429,369	В1	8/2002	Tober et al.
6,592,938	Β1	7/2003	Pessey et al.
	B2	4/2005	Mitzi
6,992,201	B2	1/2006	Scholz et al.
6,992,202	Β1	1/2006	Banger et al.
7,068,898	B2	6/2006	Buretea et al.
7,265,037	B2	9/2007	Yang et al.
7,351,282	B2	4/2008	Yamaguchi
7,466,376	B2	12/2008	Galvin et al.
7,545,051	B2	6/2009	Yang et al.
7,575,699	B2	8/2009	Strouse et al.
7,615,169	B2	11/2009	Strouse
7,892,519	B2	2/2011	Pak et al.
2002/0005145	A1	1/2002	Sherman
2002/0071970	A1	6/2002	Elder et al.
2003/0226498	A1	12/2003	Alivisatos et al.
2004/0031519	A1	2/2004	Andriessen
	A1	5/2004	Buretea et al.
2004/0120884	A1	6/2004	Sherman

2004/0126485	A1	7/2004	Thompson et al.
2004/0131934	$\mathbf{A}1$	7/2004	Sugnaux et al.
2004/0256001	A1	12/2004	Mitra et al.
2005/0016577	A1	1/2005	Andriessen et al.
2005/0133087	$\mathbf{A}1$	6/2005	Alivisatos et al.
2005/0183767	A1	8/2005	Yu et al.
2005/0267345	A1	12/2005	Korgel et al.
2005/0271827	A1	12/2005	Krunks et al.
2006/0110314	A1	5/2006	Torardi
2006/0110315	A1	5/2006	Torardi
2006/0110316	A1	5/2006	Torardi
2006/0110317	A1	5/2006	Torardi
2006/0110318	A1	5/2006	Torardi
2006/0144793	$\mathbf{A}1$	7/2006	Dadachov
2006/0159611	A1	7/2006	Hummelen et al.
2006/0216610	A1	9/2006	Galvin et al.
2006/0263291	A1	11/2006	Torardi
2007/0000537	A1	1/2007	Leidholm et al.
2007/0025139	A1	2/2007	Parsons
2007/0102040	$\mathbf{A}1$	5/2007	Beckenbaugh et al.
		(Con	tinued)
		(COII	illiaca)

FOREIGN PATENT DOCUMENTS

GB 2472541 B 3/2011

OTHER PUBLICATIONS

Deivaraj, T. et al. "Single-source precursors to ternary silver indium sulfide materials," Chem. Commun. (2001) 2304-2305.*

Deivaraj, T. et al. "Novel bimetallic thiocarboxylate compounds as single-source precursors to binary and ternary metal sulfide materials," Chem. Mater. (2003) 15: 2383-2391.*

Vittal, J. et al. "Chemistry of metal thio- and selenocarboxylates: precursors for metal sulfide/selenide materials, thin films, and nanocrystals," Acc. Chem. Res. (2006) 39: 869-877.*

Malik, M. et al. "A novel route for the preparation of CuSe and CuInSe2 nanoparticles," Advanced Materials, (1999) 11:1441-1444 *

Banger, K. et al. "Ternary single-source precursors for polycrystal-line thin-film solar cells," Appl. Organomet. Chem. (2002) 16: 617-

PCT International Search Report and Written Opinion of the International Searching Authority for PCT/US10/60583, dated Mar. 21, 2011, 12 pages.

(Continued)

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(57) ABSTRACT

Methods of forming single source precursors (SSPs) include forming intermediate products having the empirical formula $\frac{1}{2}\{L_2N(\mu-X)_2M'X_2\}_2$, and reacting MER with the intermediate products to form SSPs of the formula $L_2N(\mu-ER)_2M'$ (ER)2, wherein L is a Lewis base, M is a Group IA atom, N is a Group IB atom, M' is a Group IIB atom, each E is a Group VIB atom, each X is a Group VIIA atom or a nitrate group, and each R group is an alkyl, aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl, silane, or carbamato group. Methods of forming polymeric or copolymeric SSPs include reacting at least one of $HE^1R^1E^1H$ and MER with one or more substances having the empirical formula $L_2N(\mu-ER)_2M'(ER)_2$ or $L_2N(\mu-X)_2M'(X)_2$ to form a polymeric or copolymeric SSP. New SSPs and intermediate products are formed by such methods.

11 Claims, 3 Drawing Sheets

U.S. PATENT DOCUMENTS

20	007/0204904	A 1	9/2007	Brooks et al.
20	07/0209700	A1	9/2007	Yonezawa et al.
20	007/0277871	A1	12/2007	Lee et al.
20	07/0295385	A1	12/2007	Sheats et al.
20	008/0006322	A1	1/2008	Wang et al.
20	008/0006324	A1	1/2008	Berke et al.
20	008/0012015	A1	1/2008	Shim et al.
20	008/0023677	A1	1/2008	Frechet et al.
20	008/0026929	A1	1/2008	Jensen et al.
20	008/0031832	A1	2/2008	Wakefield et al.
20	008/0041447	A1	2/2008	Tseng et al.
20	008/0110494	A1	5/2008	Reddy
20	008/0142075	A1	6/2008	Reddy et al.
20	008/0149171	A1	6/2008	Lu et al.
20	008/0156371	A1	7/2008	LoCascio et al.
	008/0207581	A1	8/2008	Whiteford et al.
20	008/0230120	A1	9/2008	Reddy
	008/0289681	A1	11/2008	Adriani et al.
	008/0289682	A1	11/2008	Adriani et al.
	008/0308148	A1	12/2008	Leidholm et al.
	009/0050207	A1	2/2009	Galvin et al.
	009/0133751	A1	5/2009	Sreenivasan et al.
	009/0173371	A1	7/2009	Skoczenski et al.
20	009/0233398	A1	9/2009	Fox et al.

OTHER PUBLICATIONS

Bahnemann, D. W., "Ultrasmall Metal Oxide Particles: Preparation, Photophysical Characterization, and Photocatalytic Properties". Israel J. Chem., 1993, pp. 115-136, vol. 33.

Banger, K.K., et al., "Synthesis and Characterization of the First Liquid Ssingle-Source Precursors for the Deposition of Ternary Chalcopyrite (CuInS(2)) Thin Film Materials," Chem. Mater., 2001, pp. 3827-3829, vol. 13.

Banger, K.K., et al., "Facile modulation of single source precursors: the synthesis and characterization of single source precursors for deposition of ternary chalcopyrite materials," Thin Solid Films, 2002, pp. 390-395, vol. 403-404.

Banger, K.K., et al., "A New Facile Route for the Preparation of Single-Source Precursors for Bulk, Thin-Film, and Nanocrystallite 1-III-V1 Semiconductors," Inorg. Chem., 2003, pp. 7713-7715, vol. 42, No. 24.

Cardellicchio, N., et al., "Optimization of Microwave Digestion for Mercury Determination in Marine Biological Samples by Cold Vapour Atomic Absorption Spectrometry", Annali di Chimica, 2006, pp. 159-165, vol. 96 (3-4).

Carro, N., et al., "Microwave-assisted solvent extraction and gas chromatography ion trap mass spectrometry procedure for the determination of persistent organochlorine pesticides (POPs) in marine sediment", Anal. Bioanal. Chem., 2006, pp. 901-909, vol. 385.

Castro, S.L., et al., "Nanocrystalline Chalcopyrite Materials (CuInS(2)), and CuInSe(2)) via Low-Temperature Pyrolysis of Molecular Single-Source Precursors", Chem. Mater., 2003, pp. 3142-3147, vol. 15.

Castro, S.L., et al., "Synthesis and Characterization of Colloidal CulnS(2) Nanoparticles from a Molecular Single-Source Precursor," J Phys Chem B., 2004, pp. 12429-12435, vol. 108.

Choi, S.H., et al., "One-Pot Synthesis of Copper-Indium, Sulfide Nanocrystal Heterostructures with Acorn, Bottle, and Larva Shapes," J. Am Chem Soc., 2006, pp. 2520-2521, vol. 128 (8).

Domini, C.E., et al., "Comparison of three optimized digestion methods for rapid determination of chemical oxygen demand: Closed microwaves, open microwaves and ultrasound irradiation," Analytica Chimica Acta., 2006, pp. 210-217, vol. 561.

Dutta, D.P., et al., "A facile route to the synthesis of CulnS(2) nanoparticles," Materials Letters, 2006, pp. 2395-2398, vol. 60.

Gamallo-Lorenzo, D., et al., "Microwave-assisted alkaline digestion combined with microwave-assisted distillation for the determination of iodide and total iodine in edible seaweed by catalytic spectrophotometry," Analytica Chimica Acta., 2005, pp. 287-295, vol. 542.

Garcia-Vidal, J.A., et al., "Green chemistry: Efficient epoxides ringopening with 1-butanol under microwave irradiation," Applied Surface Science, 2006, pp. 6064-6066, vol. 252 (17). Gratzel, M., "Perspectives for Dye-sensitized Nanocrystalline Solar Cells," Progress in Photovoltaics: Research and Applications, 2000, pp. 171-185, vol. 8.

Grisaru, H, et al., "Microwave-Assisted Polyol Synthesis of CulnTe(2) and CulnSe(2) Nanoparticles," Inorg. Chem., 2003, pp. 7148-7155. No. 22, vol. 42.

Hayes, B.L., "Recent Advances in Microwave-Assisted Synthesis," Aldrichimica Acta., 2004, pp. 66-77, vol. 37 No. 2.

Huynh, W.U., et al., "CdSe Nanocrystal Rods/Poly (3-hexylthiophene) Composite Photovoltaic Devices," Advanced Materials, 1999, pp. 923-927, vol. 11 No. 11.

Huynh, W.U., et al., "Hybrid Nanorod-Polymer Solar Cells," Science, 2002, pp. 2425-2427, vol. 295.

Luque, A., et al., "Increasing the Efficiency of Ideal Solar Cells by Photon Induced Transitions at Intermediate Levels," Physical Review Letters, 1997, pp. 5014-5017, vol. 78 No. 26.

Marcinkevicius, S., et al., "Changes in carrier dynamics induced by proton irradiation in quantum dots," Physica. B, Condensed Matter, 2002, pp. 203-206, vol. 314.

Murali, A., et al., Synthesis and Characterization of Indium Oxide Nanoparticles, Nano Letters, 2001, pp. 287-289, vol. 1, No. 6.

Navarro, P., et al., "Optimisation of microwave assisted digestion of sediments and determination of Sn and Hg," Analytica Chimica Acta, 2006, pp. 37-44, vol. 566.

Nüchter, M., et al., "Microwave-Assisted Chemical Reactions," Chem. Eng. Technol., 2003, 1207-1216, vol. 26 (12).

Nüchter, M., et al., "Microwave Assisted Synthesis—a critical technology overview," Green Chem., 2004, pp. 128-141, vol. 6 (3).

Pak, J.J., et al., "An efficient synthesis of 4, 4', 5,5'-tetraiododibenzo-24-crown-8 and its highly conjugated derivatives," Tetrahedron Letters, 2006, pp. 233-237, vol. 47.

Perozo-Rondon, E., et al., "Microwave enhanced synthesis of N-propargyl derivatives of imidazole a green approach for the preparation of fungicidal compounds," Applied Surface Science, 2006, pp. 6067-6070, vol. 252 (17).

Sobolev, N.A., et al., "Enhanced Radiation Hardness of InAs/GaAs quantum Dot Structures," Phys. Stat. Sol. (B), 2001, pp. 93-96, vol. 224, No. 1.

Tang, Z., et al., "Semiconductor Nanoparticles on Solid Substrates: Film Structure, Intermolecular Interactions, and Polyelectrolyte Effects," Langmuir, 2002, pp. 7035-7740, vol. 18.

Walters, R.J., et al., "Radiation Hard Multi-quantum Well InP/InAsP Solar Cells for Space Applications," Progress in Photovoltaics: Research and Applications, 2000, pp. 349-354, vol. 8.

Wang, Y., et al., "Nanometer-Sized Semiconductor Clusters: Materials Synthesis, Quantum Size Effects, and Photophysical Properties," J. Phys. Chem., 1991, pp. 525-532, vol. 95.

Wei, Q., et al., "Synthesis of CulnS(2) Nanocubes by a Wet Chemical Process," Journal of Dispersion Science and Technology, 2005, pp. 555-558, vol. 26.

Zhang, X., et al., "Applications of microwave dielectric heating in environment-related heterogeneous gas-phase catalytic systems," Inorganica Chimica Acta, 2006, pp. 3421-3433, vol. 359.

Zhu, J., et al., "Microwave assisted preparation of CdSe, PbSe, andCU2-x Se nanoparticles," J. Phys. Chem. B., 2000, 104 (31), 7344-7347

Tomalia, D.A., "Birth of a New Macromolecular Architecture: Dendrimers as Quantized Building Blocks for Nanoscale Synthetic Organic Chemistry," Aldrichimica ACTA, 2004, pp. 39-57, vol. 27, No. 2.

Gerbec, J.A., et al., "Microwave-Enhanced Reaction Rates for Nanoparticle Synthesis," J. Am. Chem. Soc., 2005, pp. 15791-15800, vol. 127.

Kim, K, et al., "Synthesis of CulnSe(2) and CulnGaSe(2) Nanoparticles by Solvothermal Route," Materials Science Forum, 2004, pp. 273-276, vols. 449-452.

Bamba et al., "TiO2—ZnO Porous Films Formed by ZnO Dissolution," AZojomo, vol. 3, Dec. 2007, 7 pages.

Banger et al., "A review of single source precursors for the deposition of ternary chalcopyrite materials," NASA Conference Publication (2002), 17th Space Photovoltaic Research and Technology Conference, 2001, pp. 115-125.

Connor et al., "Phase Transformation of Biphasic CuS#CulnS to Monophasic CulnS Nanorods," J. Am. Chem. Soc, 2009, 131 (13), 4962-4966.

De Faria, et al., "Sol-Gel TiO2 Thin Films Sensitized with the Mulberry Pigment Cyanidin," Materials Research, vol. 10, No. 4, 413-417, 2007.

Deniozou et al., "Surface structure of CuGASe2 (001)" Thin Solid Films 480-481 (2005) 382-387.

Gardner et al., "Rapid synthesis and size control of CulnS2 semiconductor nanoparticles using microwave irradiation," J. Nanoparticle Research 2008, 10(4), pp. 633-641.

Halgand et al., "Physico-chemical characterisation of Cu(ln,Al)Se2 thin film for solar cells obtained by a selenisation process," Thin Solid Films 480-481 (2005) 443-446.

Hamid et al., "Preparation of Titanium Dioxide (TiO2) thin films by sol gel dip coating method," Malaysian Journal of Chemistry, 2003, vol. 5, No. 1, pp. 086-091.

Hirashima et al., "Preparation of meso-porous TiO2 gels and their characterization," Journal of Non-Crystalline Solids 285 (2001) pp. 96-100.

Jin et al., "Solar cells fabricated with CuInS2 films deposited using single-source precursors," Proceedings of the 19th European Photovoltaic Solar Energy Conference, 4AV.1.71, 2004.

Liu et al., "Preparation and characterization of CulnS2 thin films completely converted from CulnSe2 by sulfurization," Thin Solid Films 480-481 (2005) 50-54.

Miki et al., "Influence of calcination temperature on the microstructure of pourous TiO2 film," Materials Science Forum, vol. 569 (2008) pp. 17-20.

Nairn et al., "Preparation of Ultrafine Chalcopyrite Nanoparticles via the Photochemical Decomposition of Molecular Single-Source Precursors," Nano Letters 2006, vol.6(6), pp. 1218-1223.

Narako et al., "Synthesis of Metastable Wurtzite CulnSe2Nanocrystals," Chem. Mater. 2010, 22, 1613-1615.

Paez et al., "Properties of Sol-Gel TiO2 Layers on Glass Substrate," Ceramics—Silikáty 48 (2) pp. 66-71 (2004).

Pan et al., "Synthesis of Cu-In-S Ternary Nanocrystals with Tunable Structure and Composition," J. Am. Chem. Soc. Apr. 30, 2008; 130(17):5620-1, Epub Apr. 9, 2008.

PCT International Preliminary Report on Patentability for International Searching Authority for PCT/US09/36221, dated Sep. 14, 2010, 8 pages.

PCT International Search Report and Written Opinion of the International Searching Authority for PCT/US09/36221, dated Nov. 2, 2009, 11 pages.

Qi et al., "Synthesis and Characterization of Nanostructured Wurtzite CulnS2: A New Cation Disordered Polymorph of CulnS2," J. Phys. Chem. C 2009, 113, 3939-3944.

Sayilkan et al., "Characterization of TiO2 Synthesized in Alcohol by a Sol-Gel Process: The Effects of Annealing Temperature and Acid Catalyst," Turk J Chem, 29 (2005) pp. 697-706.

Sun et al., "A High-Yield Synthesis of Chalcopyrite CulnS2 Nanoparticles with Exceptional Size Control," Journal of Nanomaterials, vol. 2009, Article ID 748567, 7 pages.

Wang et al., "Synthesis of Monodispersed Wurtzite Structure CulnSe2 Nanocrystals and Their Application in High-Performance Organic-Inorganic Hybrid Photodetectors," J. Am. Chem. Soc., 2010, 132 (35), pp. 12218-12221.

Hirpo, Wakgari, et al., Synthesis of Mixed Copper-Indium Chalcogenolates. Single-Source Precursors for the Photovoltaic Materials CulnQ(2) (Q=S, Se), J. Am. Chem. Soc., 1993, pp. 1597-1599, vol. 115, No. 4.

Rodriguez, Rene G., et al., Pulsed-Spray Radiofrequency Plasma Enhanced Chemical Vapor Deposition of CulnS(2) Thin Films, Plasma Chemistry and Plasma Processing, Apr. 2006, pp. 137-148, vol. 26, No. 2.

Rodriguez, Rene, et al., The Formation of Copper Indium Disulfide Nano-Particles in Supercritical Carbon Dixoide, Program and Abstracts of the 62nd Northwest Regional Meeting of the American Chemical Society, Jun. 17-20, 2007, Boise, Idaho, p. 133, Abstract No. 232.

^{*} cited by examiner

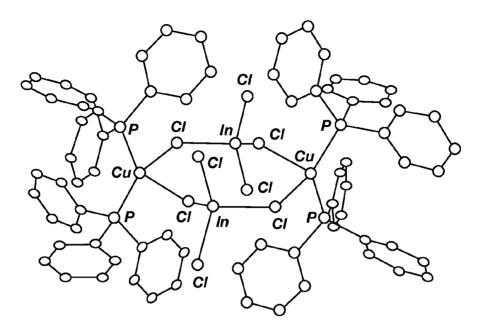


FIG. 1

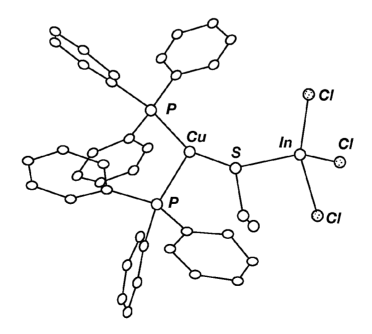


FIG. 2

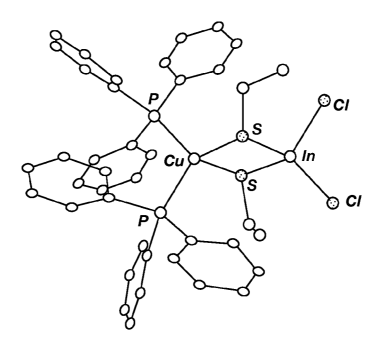


FIG. 3

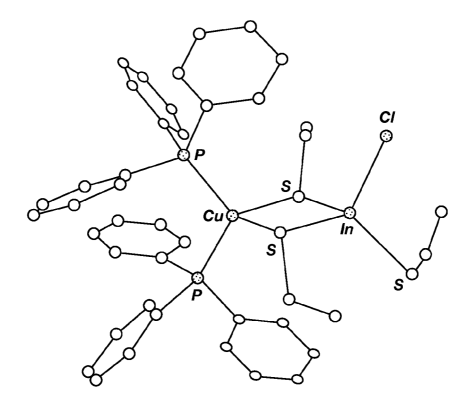


FIG. 4

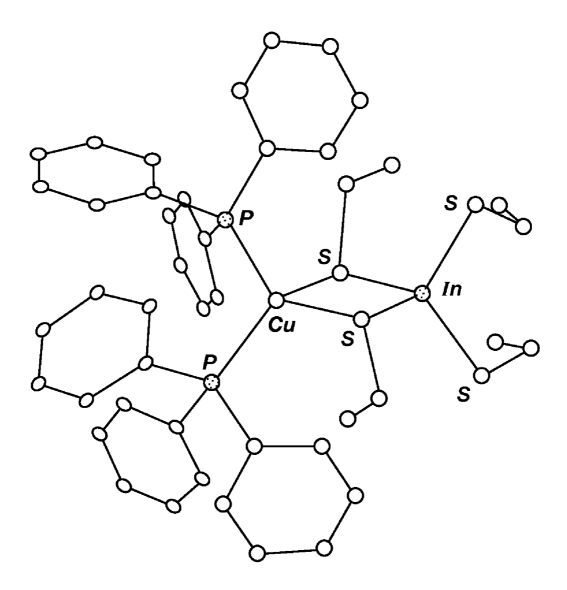


FIG. 5

METHODS OF FORMING SINGLE SOURCE PRECURSORS, METHODS OF FORMING POLYMERIC SINGLE SOURCE PRECURSORS, AND SINGLE SOURCE PRECURSORS AND INTERMEDIATE PRODUCTS FORMED BY SUCH METHODS

GOVERNMENT RIGHTS

This invention was made under a Cooperative Research and Development Agreement between Precision Nanoparticles and Battelle Energy Alliance, LLC under Contract Number DE-AC07-05ID14517 awarded by the United States Department of Energy. The government has certain rights in this invention.

CROSS-REFERENCE TO RELATED APPLICATIONS

The subject matter of this application is related to the subject matter of U.S. patent application Ser. No. 13/191,062, filed Jul. 26, 2011, to the subject matter of U.S. patent application Ser. No. 13/099,043, filed May 2, 2011, which application is a divisional of U.S. patent application Ser. No. 25 12/047,956, filed Mar. 13, 2008, now U.S. Pat. No. 8,003,070, issued Aug. 23, 2011, and to the subject matter of U.S. patent application Ser. No. 13/019,879, filed Feb. 2, 2011.

TECHNICAL FIELD

Embodiments of the invention relate to methods of forming single source precursors and, more particularly, to methods of forming organometallic single source precursors for use in forming semiconductor materials, particles, and devices.

BACKGROUND

Semiconductor devices are devices that employ semiconductor materials, which are solid materials that exhibit an 40 electrical conductivity lying between that of a conductor and that of an insulator. Semiconductor devices include, for example, diodes (e.g., light emitting diodes (LEDs)), photovoltaic devices, sensors, solid state lasers, and integrated circuits (e.g., memory modules and microprocessors).

Semiconductor materials that can be employed in semiconductor devices include, for example, silicon (Si), germanium (Ge), chalcopyrites (e.g., CuInS $_2$, CuGaS $_2$, and CuInSe $_2$), chalcogenides (e.g., Cu(In $_x$ Ga $_{1-x}$)(Se $_y$ S $_{1-y}$) $_2$), cadmium telluride (CdTe), gallium arsenide (GaAs), organic polymers (e.g., 50 polyphenylene vinylene, copper phthalocyanine, fullerenes), and light absorbing dyes (e.g., ruthenium-centered metalorganic dyes).

It has been discovered that chalcopyrite materials may be formed by decomposing one or more so-called "single source 55 precursors" (SSPs), which are organometallic substances (e.g., molecules, complexes, etc.) that comprise all of the atomic elements, in the appropriate stoichiometric ratios, necessary to form a chalcopyrite material. Such methods, and methods of forming such SSPs are disclosed in, for example, 60 Hirpo et al., *Synthesis of Mixed Copper-Indium Chalcogenolates. Single-Source Precursors for The Photovoltaic Materials CuInQ*₂ (Q=S, Se), Journal Of The American Chemical Society, Vol. 115, Iss. 4, pp. 1597-1599 (Feb. 24, 1993). Methods for forming such SSPs are also disclosed in, for example, U.S. Pat. No. 6,992,202, which issued Jan. 31, 2006 to Banger et al.

2

There remains a need in the art, however, for improved methods that may be used to form single source precursors for use in forming chalcopyrite materials (e.g., semiconductive ternary chalcopyrite materials).

BRIEF SUMMARY

In some embodiments, the present invention includes methods of forming single source precursors. The methods include forming an intermediate product having the empirical formula $^{1\!/2}\{L_2N(\mu\text{-}X)_2M'X_2\}_2$, and reacting MER with the intermediate product to form a single source precursor having the empirical formula $L_2N(\mu\text{-}ER)_2M'(ER)_2$, wherein L is a Lewis base coordinated to N by a dative bond, each M is individually selected from Group IA atoms, each M is individually selected from Group IB atoms, each M is individually selected from Group IIIA atoms, each E is individually selected from Group VIA atoms, each X is individually selected from Group VIIA atoms or a nitrate group, and each R group is individually selected from the group consisting of alkyl, aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl, silane, and carbamato groups.

In some embodiments, the present invention includes methods of forming single source precursors from other single source precursors. For example, HE^2R^2 may be reacted with a first single source precursor having the empirical formula $L_2N(\mu-E^1R^1)_2M'(E^1R^1)_2$ to form a second, different single source precursor having the empirical formula $L_2N(\mu-E^2R^2)_2M'(E^2R^2)_2$, wherein L is a Lewis base coordinated to N by a dative bond, each N is individually selected from Group IIB atoms, each M' is individually selected from Group VIA atoms, and each R^1 and R^2 group is individually selected from the group consisting of alkyl, aryl, vinyl, (per)fluoro 35 alkyl, (per)fluoro aryl, silane, and carbamato groups.

In additional embodiments, the present invention includes methods of forming polymeric single source precursors, in which HE¹R¹E¹H is reacted with a single source precursor having the empirical formula L₂N(µ-ER)₂M'(ER)₂ to form a polymeric single source precursor having the empirical for- $[L_2N(\mu-ER)_a(\mu-E^1R^1E^1)_bM'(ER)_c(E^1R^1E^1)_d]_m$ mula wherein L is a Lewis base coordinated to N by a dative bond, each N is individually selected from Group IB atoms, each M' is individually selected from Group IIIA atoms, each E and E¹ is individually selected from Group VIA atoms, a is any number from zero (0) to two (2), b is the difference between two (2) and a (b=2-a), c is any number from zero (0) to two (2), d is the difference between two (2) and c (d=2-c), m is any number, each of the R groups is individually selected from the group consisting of alkyl, aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl, silane, and carbamato groups, and each of the R¹ groups is individually selected from the group consisting of aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl, silane, carbamato, and α, ω -E' functionalized alkyl groups.

In additional embodiments, the present invention includes methods of forming polymeric single source precursors in which at least one of $HE^1R^1E^1H$ and HE^2R^2 is reacted with a single source precursor having the empirical formula $L_2N(\mu-ER)_2M'(ER)_2$ to form a polymeric single source precursor having the empirical formula $[L_2N(E^2R^2)_a(\mu-E^1R^1E^1)_bM'(E^2R^2)_c(E^1R^1E^1)_a]_m$, wherein L is a Lewis base coordinated to N by a dative bond, each N is individually selected from Group IB atoms, each M' is individually selected from Group IIA atoms, each E, E^1 , and E^2 is individually selected from Group VIA atoms, a is any number from zero (0) to two (2), b is the difference between two (2) and a, c is any number from zero (0) to two (2), d is the difference between two (2)

and c, m is any number, each R and R² is individually selected from the group consisting of alkyl, aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl, silane, and carbamato groups, and each R¹ is individually selected from the group consisting of aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl, silane, carba-5 mato, and α,ω -E' functionalized alkyl groups.

In yet further embodiments, the present invention includes methods of forming copolymeric single source precursors in which a first single source precursor, a second single source precursor differing from the first single source precursor, and 10 HE¹R¹E¹H are reacted to form a copolymeric single source precursor having the empirical formula $\{[L_2N^1(ER)_a(\mu [E^{1}R^{1}E^{1})_{b}M^{1}(ER)_{c}(E^{1}R^{1}E^{1})_{d}]_{m}[L_{2}N^{2}(ER)_{e}(\mu-E^{1}R^{1}E^{1})_{c}M^{2}]$ $(ER)_g(E^1R^1E^1)_h]_n\}_l$, wherein L is a Lewis base, each N^1 and N² is individually selected from Group IB atoms, each M'¹ and M² is individually selected from Group IIIA atoms, each E and E¹ is individually selected from Group VIA atoms, a is any number from zero (0) to two (2), b is the difference between two (2) and a, c is any number from zero (0) to two (2), d is the difference between two (2) and c, e is any number 20 from zero (0) to two (2), f is the difference between two (2) and e, g is any number from zero (0) to two (2), h is the difference between two (2) and g, m is any number, n is any number, 1 is any number, each R is individually selected from the group consisting of alkyl, aryl, vinyl, (per)fluoro alkyl, 25 (per)fluoro aryl, silane, and carbamato groups, and each R¹ is individually selected from the group consisting of aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl, silane, carbamato, and α, ω -E¹ functionalized alkyl groups.

Additional embodiments of the present invention include 30 methods of forming polymeric single source precursors in which at least one of ME¹R¹E¹M and MER is reacted with a substance having the empirical formula $L_2N(\mu-X)_2M'(X)_2$ to form a polymeric single source precursor having the empirical formula $[L_2N(\mu-ER)_a(\mu-E^1R^1E^1)_bM'(ER)_c(E^1R^1E^1)_d]_m$, 35 wherein L is a Lewis base that is coordinated to N by a dative bond, each M is individually selected from Group IA atoms, each N is individually selected from Group IB atoms, each M' is individually selected from Group IIIA atoms, each E and E¹ is individually selected from Group VIA atoms, each X is 40 individually selected from Group VIIA atoms or a nitrate group, a is any number from zero (0) to two (2), b is the difference between two (2) and a (b=2-a), c is any number from zero (0) to two (2), d is the difference between two (2) and c (d=2-c), m is any number, each of the R groups is 45 individually selected from the group consisting of alkyl, aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl, silane, and carbamato groups, and each of the R¹ groups is individually selected from the group consisting of aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl, silane, carbamato, and α,ω -E¹ func- 50 tionalized alkyl groups.

In additional embodiments, copolymeric single source precursors are formed by reacting uME¹R¹E¹M and vMER with a first substance having the empirical formula $L_2N^1(\mu-X)_2M^{-1}$ (X)₂ and a second substance having the empirical formula 55 $L_2N^2(\mu-X)_2M^2(X)_2$ to form a copolymeric single source precursor having the empirical formula $\{[L_2N^1(ER)_a(\mu E^{1}R^{1}E^{1})_{b}M^{1}(ER)_{c}(E^{1}R^{1}E^{1})_{d}]_{m}[L_{2}N^{2}(ER)_{e}(\mu-E^{1}R^{1}E^{1})_{f}M^{2}]$ $(ER)_{\mathfrak{g}}(E^1R^1E)_h]_n$, wherein L is a Lewis base, each N¹ and N² is individually selected from Group IB atoms, each M¹ and M¹² is individually selected from Group IIIA atoms, each E and E¹ is individually selected from Group VIA atoms, a is any number from zero (0) to two (2), b is the difference between two (2) and a, c is any number from zero (0) to two from zero (0) to two (2), f is the difference between two (2) and e, g is any number from zero (0) to two (2), h is the

difference between two (2) and g, m is any number, n is any number, l is any number, u is any number from zero (0) to four (4), v is the difference between four (4) and u, each R is individually selected from the group consisting of alkyl, aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl, silane, and carbamato groups, and each R¹ is individually selected from the group consisting of aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl, silane, carbamato, and α,ω -E¹ functionalized alkyl groups.

Further embodiments of the present invention include single source precursors, polymeric single source precursors, and copolymeric single source precursors formed by the methods described herein.

For example, in some embodiments, the present invention includes polymeric, organometallic single source precursor having the empirical formula $[L_2N(ER)_a(\mu-E^1R^1E^1)_bM']$ $(ER)_c (E^1R^1E^1)_d$ _m, wherein L is a Lewis base, each N is individually selected from Group IB atoms, each M' is individually selected from Group IIIA atoms, each E and E¹ is individually selected from Group VIA atoms, a is any number from zero (0) to two (2), b is the difference between two (2) and a, c is any number from zero (0) to two (2), d is the difference between two (2) and c, m is any number, each R is individually selected from the group consisting of alkyl, aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl, silane, and carbamato groups, and each R¹ is individually selected from the group consisting of aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl, silane, carbamato, and α,ω -E¹ functionalized alkyl groups.

Yet further embodiments of the present invention include polymeric, organometallic single source precursors having the empirical formula $[L_2N(\mu ER^1)_a(\mu - ER^2E)_bM'(ER^1)_c$ $(ER^2E)_{alm}$, wherein L is a Lewis base, each N is individually selected from Group IB atoms, each M' is individually selected from Group IIIA atoms, each E is individually selected from Group VIA atoms, a is any number from zero (0) to two (2), b is the difference between two (2) and a (b=2-a), c is any number from zero (0) to two (2), d is the difference between two (2) and c (d=2-c), m is any number, each of the R¹ groups is individually selected from the group consisting of alkyl, aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl, silane, and carbamato groups, and each of the R² groups is individually selected from the group consisting of alkyl, vinyl, and aryl groups.

BRIEF DESCRIPTION OF THE SEVERAL VIEWS OF THE DRAWINGS

While the specification concludes with claims particularly pointing out and distinctly claiming that which is regarded as the present invention, advantages of this invention may be more readily ascertained from the following description of the invention when read in conjunction with the accompanying drawings in which:

FIGS. 1 through 4 illustrate various intermediate products that may be formed in accordance with embodiments of methods of the present invention for forming single source precursors; and

FIG. 5 illustrates one example of a single source precursor 60 that may be formed using methods of the present invention.

DETAILED DESCRIPTION

The illustrations presented herein are not meant to be (2), d is the difference between two (2) and c, e is any number 65 actual views of any particular molecule or complex, but are merely idealized representations that are employed to describe various embodiments of the present invention.

The chalcogens are the elements in Group VIA (16) of the periodic table. As used herein, the term "chalcogenide" means and includes any compound that contains at least one chalcogen element other than oxygen (e.g., sulfur, selenium, tellurium, etc.). Chalcogenides include, for example, ternary chalcopyrite materials.

As used herein the term "ternary chalcopyrite material" means and includes any material having a composition generally represented by the formula I-III-VI2, where roman numeral I refers to elements in Group I (Groups IA (1) and IB (11)) of the periodic table, roman numeral III refers to elements in Group III (Groups IIIB (3) and IIIA (13)) of the periodic table, and roman numeral VI refers to elements in Group VI (Groups VIB (6) and VIA (16)) of the periodic 15 table. By ternary, it is meant that the chalcopyrite materials contain atoms from three elemental Groups of the periodic table. For example, approximately twenty-five percent (25%) of the atoms in a ternary chalcopyrite material may be from Group IB, approximately twenty-five percent (25%) of the 20 [bis(4-trifluoromethyl-thiophenolato)indium]bis[µ-(4-triflatoms may be from Group IIIA, and approximately fifty percent (50%) of the atoms may be from Group VIA. CuInS₂, CuInSe₂, Cu(In,Ga)Se₂, CuGaSe₂, and AgInS₂ are examples of ternary chalcopyrite materials. It should be noted that ternary chalcopyrites include materials having multiple and/ 25 or different atoms from each of three Groups of the periodic table. For example, CuInSSe is a ternary chalcopyrite because it has Cu (Group IB), In (Group IIIA), and S and Se (both from Group VIA). In addition, materials of the form (Cu:Ag)(In: Ga)(S:Se), having various ratios of the respectively grouped atoms are all ternary chalcopyrites (Cu and Ag both are in Group IB, In and Ga both are in Group IIIA, S and Se both are in Group VIA).

As used herein, the terms "single source precursor" and "SSP" mean and include any substance (e.g., a molecule or 35 complex) that comprises all of the necessary atomic elements, in the appropriate stoichiometric ratios, necessary to form a chalcogenide material (e.g., a ternary chalcopyrite material). Single source precursors may comprise so-called organometallic substances. As non-limiting examples, single source 40 precursors include molecules or complexes having the empirical formula $[L_2N(\mu-ER)_2M(ER)_2]$ (or $L_2NM(ER)_4$), wherein L is a Lewis base that is coordinated to N by a dative bond, each N is individually selected from Group IB atoms, each M is individually selected from Group IIIA atoms, each 45 E is individually selected from Group VIA atoms, and each R is individually selected from the group consisting of alkyl. aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl, silane, and carbamato groups. As seven particular non-limiting examples, single source precursors include (Ph₃P)₂Cu(μ-SEt)₂In(SEt)₂, 50 $(Ph_3P)_2Cu(\mu-SEt)_2Ga(SEt)_2$, $(Ph_3P)_2Cu(\mu-SEt)_2Al(SEt)_2$, $(Ph_3P)_2Ag(\mu-SEt)_2In(SEt)_2$, $(Ph_3P)_2Ag(\mu-SEt)_2Ga(SEt)_2$, $(Ph_3P)_2Ag(\mu-SEt)_2Al(SEt)_2$, and $\{(Ph_3P)_2Cu(\mu-Cl)_2InCl_2\}_2$.

By way of example and not limitation, the following are examples of copper-indium SSPs:

[bis(ethanethiolato)indium]bis[μ-(ethanethiolato)]bis(triisobutylphosphine)-copper;

[bis(ethanethiolato)indium]bis[μ-(ethanethiolato)]bis(trihexylphosphine)-copper;

[bis(ethanethiolato)indium]bis[μ-(ethanethiolato)]bis(triphenylphosphine)-copper;

[bis(ethanethiolato)indium]bis[μ-(ethanethiolato)]bis(perfluorotriphenylphosphine)-copper;

[bis(propanethiolato)indium]bis[μ-(propanethiolato)]bis(triisobutylphosphine)-copper;

[bis(propanethiolato)indium]bis[μ-(propanethiolato)]bis(trihexylphosphine)-copper;

6

[bis(propanethiolato)indium]bis[\(\mu\)-(propanethiolato)]bis (triphenylphosphine)-copper;

[bis(propanethiolato)indium]bis[μ-(propanethiolato)]bis (perfluorotriphenylphosphine)-copper;

[bis(hexanethiolato)indium]bis[μ-(propanethiolato)]bis(triisobutylphosphine)-copper:

[bis(hexanethiolato)indium]bis[\u03c4-(propanethiolato)]bis(trihexylphosphine)-copper;

[bis(hexanethiolato)indium]bis[\u03c4-(propanethiolato)]bis (triphenylphosphine)-copper;

[bis(hexanethiolato)indium]bis[μ-(propanethiolato)]bis(perfluorotriphenylphosphine)-copper;

[bis(4-trifluoromethyl-thiophenolato)indium]bis[μ -(4-trifluoromethylthiophenolato)]bis(triisobutylphosphine)-

[bis(4-trifluoromethyl-thiophenolato)indium]bis[μ-(4-trifluoromethyl-thiophenolato)]bis(trihexylphosphine)-cop-

luoromethyl-thiophenolato)]bis(triphenylphosphine)copper;

[bis(4-trifluoromethyl-thiophenolato)indium]bis[\mu-(4-trifluoromethyl-thiophenolato)]bis(perfluorotriphenylphosphine)-copper;

[bis(3,5-bis(trifluoromethyl)-thiophenolato)indium]bis[μ-(3,5-bis(trifluoromethyl)-thiophenolato)]bis(triisobu-[bis(3,5-bis(trifluoromethyl)tylphosphine)-copper; thiophenolato)indium]bis[\mu-(3,5-bis(trifluoromethyl)thiophenolato)]bis(trihexylphosphine)-copper;

[bis(3,5-bis(trifluoromethyl)-thiophenolato)indium]bis[µ-(3,5-bis(trifluoromethyl)-thiophenolato)]bis(triphenylphosphine)-copper;

[bis(3,5-bis(trifluoromethyl)-thiophenolato)indium]bis[µ-(3,5-bis(trifluoromethyl)-thiophenolato)]bis(perfluorotriphenylphosphine)-copper;

[bis(thiophenolato)indium]bis[μ-(thiophenolato)]bis(triisobutylphosphine)-copper;

[bis(thiophenolato)indium]bis[μ-(thiophenolato)]bis(trihexylphosphine)-copper;

[bis(thiophenolato)indium]bis[\mu-(thiophenolato)]bis(triphenylphosphine)-copper;

[bis(thiophenolato)indium]bis[\u03c4-(thiophenolato)]bis(perfluorotriphenylphosphine)-copper;

[bis(α -toluenethiolato)indium]bis[μ -(α -toluenethiolato)]bis (triisobutylphosphine)-copper;

[bis(α -toluenethiolato)indium]bis[μ -(α -toluenethiolato)]bis (trihexylphosphine)-copper;

 $[bis(\alpha-toluenethiolato)indium]bis[\mu-(\alpha-toluenethiolato)]bis$ (triphenylphosphine)-copper;

[bis(α -toluenethiolato)indium]bis[μ -(α -toluenethiolato)]bis (perfluorotriphenylphosphine)-copper;

[bis(pentafluorothiophenolato)indium]bis[μ-(pentafluorothiophenolato)]bis(triisobutylphosphine)-copper; (pentafluorothiophenolato)indium]bis[μ-(pentafluorothiophenolato)]bis(trihexylphosphine)-copper;

[bis(pentafluorothiophenolato)indium]bis[μ-(pentafluorothiophenolato)]bis(triphenylphosphine)-copper; [bis (pentafluorothiophenolato)indium]bis[μ-(pentafluorothiophenolato)]bis(perfluorotriphenylphosphine)copper;

[bis(thiobenzoato)indium]bis[μ-(thiobenzoato)]bis(triisobutylphosphine)-copper;

[bis(thiobenzoato)indium]bis[\u03c4-(thiobenzoato)]bis(trihexylphosphine)-copper;

[bis(thiobenzoato)indium]bis[\u03c4-(thiobenzoato)]bis(triphenylphosphine)-copper;

[bis(thiobenzoato)indium]bis[\mu-(thiobenzoato)]bis(perfluorotriphenylphosphine)-copper;

[bis(thiobenzoato)indium]bis[\u03c4-(thiobenzoato)]bis[ethylenebis(diphenylphosphine)]-copper;

[bis(thiobenzoato)indium]bis[\mu-(thiobenzoato)]bis[bis(2diphenylphosphinophenyl)ether]-copper;

[bis(thiobenzoato)indium]bis[µ-(thiobenzoato)]bis(trimethylphosphine)-copper;

[bis(ethanethiolato)indium]bis[µ-(ethanethiolato)]bis[ethylenebis(diphenylphosphine)]-copper;

[bis(ethanethiolato)indium]bis[\u03c4-(ethanethiolato)]bis[bis(2diphenylphosphinophenyl)ether]-copper;

 $[bis(ethanethiolato)indium] bis[\mu\text{-}(thiophenolato)] bis(triph$ enylphosphine)-copper; and

[bis(ethanethiolato)indium]bis[\u03c4-(ethanethiolato)]bis(trim-

ethylphosphine)-copper. By way of example and not limitation, the following are examples of copper-gallium SSPs: [bis(ethanethiolato)gallium]bis[μ-(ethanethiolato)]bis(triisobutylphosphine)-copper; [bis(ethanethiolato)gallium]bis[u-(ethanethiolato)]bis 20 (trihexylphosphine)-copper; [bis(ethanethiolato)gallium]bis [μ-(ethanethiolato)]bis(triphenylphosphine)-copper; $(ethanethiolato) gallium] bis [\mu-(ethanethiolato)] bis$ (perfluorotriphenylphosphine)-copper; [bis(propanethiolato) $gallium] bis [\mu\text{-(propanethiolato)}] bis (triis obutyl phosphine)\text{-}$ [bis(propanethiolato)gallium]bis[μ-(propanethiolato)]bis(trihexylphosphine)-copper; **Fbis** (propanethiolato)gallium]bis[μ-(propanethiolato)]bis (triphenylphosphine)-copper; [bis(propanethiolato)gallium] bis[μ-(propanethiolato)]bis(perfluorotriphenylphosphine)copper; [bis(hexanethiolato)gallium]bis[μ-(propanethiolato)]bis(triisobutylphosphine)-copper; [bis $(hexanethiolato)gallium] bis [\mu-(propanethiolato)] bis$ (trihexylphosphine)-copper; [bis(hexanethiolato)gallium]bis [μ-(propanethiolato)]bis(triphenylphosphine)-copper; [bis 35 (hexanethiolato)gallium]bis[μ-(propanethiolato)]bis (perfluorotriphenylphosphine)-copper; [bis(4trifluoromethyl-thiophenolato)gallium]bis[μ-(4trifluoromethyl-thiophenolato)]bis(triisobutylphosphine)copper; [bis(4-trifluoromethyl-thiophenolato)gallium]bis[μ- 40 (4-trifluoromethyl-thiophenolato)]bis(trihexylphosphine)copper; [bis(4-trifluoromethyl-thiophenolato)gallium]bis[μ-(4-trifluoromethyl-thiophenolato)]bis(triphenylphosphine)copper; [bis(4-trifluoromethyl-thiophenolato)gallium]bis[μ-(4-trifluoromethyl-thiophenolato)]bis (perfluorotriphenylphosphine)-copper; [bis(3,5-bis (trifluoromethyl)-thiophenolato)gallium]bis[μ -(3,5-bis (trifluoromethyl)-thiophenolato)]bis(triisobutylphosphine)-[bis(3,5-bis(trifluoromethyl)-thiophenolato) gallium]bis[μ -(3,5-bis(trifluoromethyl)-thiophenolato bis 50 (trihexylphosphine)-copper; [bis(3,5-bis(trifluoromethyl)thiophenolato)gallium]bis[μ-(3,5-bis(trifluoromethyl)thiophenolato)]bis(triphenylphosphine)-copper; [bis(3,5-bis (trifluoromethyl)-thiophenolato)gallium]bis[μ -(3,5-bis (trifluoromethyl)-thiophenolato)]bis (perfluorotriphenylphosphine)-copper; [bis(thiophenolato) gallium]bis[u-(thiophenolato)]bis(triisobutylphosphine)copper; [bis(thiophenolato)gallium]bis[μ-(thiophenolato)] bis(trihexylphosphine)-copper; [bis(thiophenolato)gallium] bis[μ-(thiophenolato)]bis(triphenylphosphine)-copper; [bis 60 (thiophenolato)gallium]bis[u-(thiophenolato)]bis (perfluorotriphenylphosphine)-copper; [bis(α-

toluenethiolato)gallium]bis[μ -(α -toluenethiolato)]bis

toluenethiolato)]bis(triphenylphosphine)-copper;

gallium]bis[μ -(α -toluenethiolato)]bis(trihexylphosphine)-

(triisobutylphosphine)-copper;

[bis(α -toluenethiolato)

[bis(α-

[bis(α -toluenethiolato)gallium]bis[μ -(α -

8

toluenethiolato)gallium]bis[μ -(α -toluenethiolato)]bis(perfluorotriphenylphosphine)-copper; [bis (pentafluorothiophenolato)gallium]bis[μ-(pentafluorothiophenolato)]bis(triisobutylphosphine)-[bis(pentafluorothiophenolato)gallium]bis[μcopper; (pentafluorothiophenolato)]bis(trihexylphosphine)-copper; [bis(pentafluorothiophenolato)gallium]bis[µ-(pentafluorothiophenolato)]bis(triphenylphosphine)-copper; and [bis (pentafluorothiophenolato)gallium]bis[μ-(pentafluo-

rothiophenolato)]bis(perfluorotriphenylphosphine)-copper. By way of example and not limitation, the following are examples of silver-indium SSPs: [bis(ethanethiolato)indium] $bis[\mu-(ethanethiolato)]bis(triisobutylphosphine)-silver;$ [bis $(ethanethiolato) indium] bis [\mu-(ethanethiolato)] bis (trihexy-$ [bis(ethanethiolato)indium]bis[μlphosphine)-silver; (ethanethiolato)]bis(triphenylphosphine)-silver; (ethanethiolato)indium]bis[μ-(ethanethiolato)]bis (perfluorotriphenylphosphine)-silver; [bis(propanethiolato) indium]bis[μ-(propanethiolato)]bis(triisobutylphosphine)silver: [bis(propanethiolato)indium]bis[u-(propanethiolato)] bis(trihexylphosphine)-silver; [bis(propanethiolato)indium] bis[μ-(propanethiolato)]bis(triphenylphosphine)-silver; [bis $(propanethiolato)indium]bis[\mu-(propanethiolato)]bis$ (perfluorotriphenylphosphine)-silver; [bis(hexanethiolato) indium]bis[\(\mu\)-(propanethiolato)]bis(triisobutylphosphine)silver; [bis(hexanethiolato)indium]bis[μ-(propanethiolato)] bis(trihexylphosphine)-silver; [bis(hexanethiolato)indium] bis[μ-(propanethiolato)]bis(triphenylphosphine)-silver; [bis (hexanethiolato)indium]bis[μ-(propanethiolato)]bis (perfluorotriphenylphosphine)-silver; [bis(4trifluoromethyl-thiophenolato)indium]bis[μ-(4trifluoromethyl-thiophenolato)]bis(triisobutylphosphine)-[bis(4-trifluoromethyl-thiophenolato)indium]bis[μ-(4-trifluoromethyl-thiophenolato)]bis(trihexylphosphine)silver; [bis(4-trifluoromethyl-thiophenolato)indium]bis[μ-(4-trifluoromethyl-thiophenolato)]bis(triphenylphosphine)silver; [bis(4-trifluoromethyl-thiophenolato)indium]bis[µ-(4-trifluoromethyl-thiophenolato)]bis (perfluorotriphenylphosphine)-silver; [bis(3,5-bis(trifluoromethyl)-thiophenolato)indium]bis[\mu-(3,5-bis (trifluoromethyl)-thiophenolato)]bis(triisobutylphosphine)silver; [bis(3,5-bis(trifluoromethyl)-thiophenolato)indium] bis[μ-(3,5-bis(trifluoromethyl)-thiophenolato)]bis (trihexylphosphine)-silver; [bis(3,5-bis(trifluoromethyl)thiophenolato)indium]bis[μ-(3,5-bis(trifluoromethyl)thiophenolato)]bis(triphenylphosphine)-silver; and [bis(3,5bis(trifluoromethyl)-thiophenolato)indium]bis[u-(3,5-bis (trifluoromethyl)-thiophenolato)]bis (perfluorotriphenylphosphine)-silver.

By way of example and not limitation, the following are examples of silver-gallium SSPs: [bis(ethanethiolato)gallium]bis[u-(ethanethiolato)]bis(triisobutylphosphine)-silver; [bis(ethanethiolato)gallium]bis[μ-(ethanethiolato)]bis (trihexylphosphine)-silver; [bis(ethanethiolato)gallium]bis [μ-(ethanethiolato)]bis(triphenylphosphine)-silver; (ethanethiolato)gallium]bis[μ-(ethanethiolato)]bis (perfluorotriphenylphosphine)-silver; [bis(propanethiolato) $gallium] bis [\mu-(propanethiolato)] bis (triis obutyl phosphine)$ silver; [bis(propanethiolato)gallium]bis[µ-(propanethiolato)]bis(trihexylphosphine)-silver; (propanethiolato)gallium]bis[μ-(propanethiolato)]bis (triphenylphosphine)-silver; [bis(propanethiolato)gallium] bis[μ-(propanethiolato)]bis (perfluorotriphenylphosphine)silver; [bis(hexanethiolato)gallium]bis[μ-(propanethiolato)] bis(triisobutylphosphine)-silver; [bis(hexanethiolato) gallium]bis[μ-(propanethiolato)]bis(trihexylphosphine)silver; [bis(hexanethiolato)gallium]bis[μ-(propanethiolato)]

bis(triphenylphosphine)-silver; [bis(hexanethiolato)gallium] bis[μ-(propanethiolato)]bis(perfluorotriphenylphosphine)silver; [bis(4-trifluoromethyl-thiophenolato)gallium]bis[μ-(4-trifluoromethyl-thiophenolato)lbis [bis(4-trifluoromethyl-5 (triisobutylphosphine)-silver; thiophenolato)gallium]bis[u-(4-trifluoromethylthiophenolato) lbis(trihexylphosphine)-silver: Tbis(4trifluoromethyl-thiophenolato)gallium]bis[μ-(4trifluoromethyl-thiophenolato)]bis(triphenylphosphine)silver; [bis(4-trifluoromethyl-thiophenolato)gallium]bis[μ-(4-trifluoromethyl-thiophenolato)]bis (perfluorotriphenylphosphine)-silver; [bis(3,5-bis (trifluoromethyl)-thiophenolato)gallium]bis[μ-(3,5-bis (trifluoromethyl)-thiophenolato)]bis(triisobutylphosphine)silver; [bis(3,5-bis(trifluoromethyl)-thiophenolato)gallium] bis $[\mu$ -(3,5-bis(trifluoromethyl)-thiophenolato)] bis (trihexylphosphine)-silver; [bis(3,5-bis(trifluoromethyl)thiophenolato)gallium]bis[μ-(3,5-bis(trifluoromethyl)thiophenolato)]bis(triphenylphosphine)-silver; [bis(3,5-bis 20 [bis(3,5-bis(trifluoromethyl)-thiophenolato)aluminum]bis (trifluoromethyl)-thiophenolato)gallium]bis[μ -(3,5-bis (trifluoromethyl)-thiophenolato)]bis (perfluorotriphenylphosphine)-silver; [bis(thiophenolato) gallium]bis[μ-(thiophenolato)]bis(triisobutylphosphine)silver; [bis(thiophenolato)gallium]bis[μ-(thiophenolato)]bis 25 (trihexylphosphine)-silver; [bis(thiophenolato)gallium]bis [μ-(thiophenolato)]bis(triphenylphosphine)-silver; [bis (thiophenolato)gallium]bis[μ-(thiophenolato)]bis (perfluorotriphenylphosphine)-silver; [bis(αtoluenethiolato)gallium]bis[μ -(α -toluenethiolato)]bis (triisobutylphosphine)-silver; [bis(α -toluenethiolato) gallium]bis[μ -(α -toluenethiolato)]bis(trihexylphosphine)-[bis(α -toluenethiolato)gallium]bis[μ -(α silver: toluenethiolato)]bis(triphenylphosphine)-silver; [bis(αtoluenethiolato)gallium]bis[μ -(α -toluenethiolato)]bis (perfluorotriphenylphosphine)-silver; This (pentafluorothiophenolato)gallium]bis[u-(pentafluorothiophenolato)]bis(triisobutylphosphine)-silver; [bis(pentafluorothiophenolato)gallium]bis[µ-(pentafluorothiophenolato)]bis(trihexylphosphine)-silver; [bis(pen- 40 tafluorothiophenolato)gallium]bis[μ-(pentafluorothiophenolato)]bis(triphenylphosphine)-silver; bis

(perfluorotriphenylphosphine)-silver. By way of example and not limitation, the following are examples of copper-aluminum SSPs:

(pentafluorothiophenolato)gallium]bis[µ-

(pentafluorothiophenolato)]bis

[bis(ethanethiolato)aluminum]bis[µ-(ethanethiolato)]bis(triisobutylphosphine)-copper;

[bis(ethanethiolato)aluminum]bis[\mu-(ethanethiolato)]bis(tri- 50 hexylphosphine)-copper;

[bis(ethanethiolato)aluminum]bis[u-(ethanethiolato)]bis (triphenylphosphine)-copper;

[bis(ethanethiolato)aluminum]bis[μ-(ethanethiolato)]bis (perfluorotriphenylphosphine)-copper;

[bis(propanethiolato)aluminum]bis[\u03c4-(propanethiolato)]bis (triisobutylphosphine)-copper;

[bis(propanethiolato)aluminum]bis[μ-(propanethiolato)]bis (trihexylphosphine)-copper;

[bis(propanethiolato)aluminum]bis[u-(propanethiolato)]bis (triphenylphosphine)-copper;

[bis(propanethiolato)aluminum]bis[μ-(propanethiolato)]bis (perfluorotriphenylphosphine)-copper; [bis(hexanethiolato)aluminum]bis[μ-(propanethiolato)]bis(triisobutylphosphine)-copper;

[bis(hexanethiolato)aluminum]bis[µ-(propanethiolato)]bis (trihexylphosphine)-copper;

10

[bis(hexanethiolato)aluminum]bis[μ-(propanethiolato)]bis (triphenylphosphine)-copper;

[bis(hexanethiolato)aluminum]bis[μ-(propanethiolato)]bis (perfluorotriphenylphosphine)-copper;

[bis(4-trifluoromethyl-thiophenolato)aluminum]bis[\u03c4-(4trifluoromethylthiophenolato)]bis(triisobutylphosphine)-

[bis(4-trifluoromethyl-thiophenolato)aluminum]bis[\u03c4-(4trifluoromethyl-thiophenolato)]bis(trihexylphosphine)-

[bis(4-trifluoromethyl-thiophenolato)aluminum]bis[\u03c4-(4trifluoromethyl-thiophenolato)]bis(triphenylphosphine)copper;

[bis(4-trifluoromethyl-thiophenolato)aluminum]bis[μ-(4trifluoromethyl-thiophenolato)]bis(perfluorotriphenylphosphine)-copper;

[bis(3,5-bis(trifluoromethyl)-thiophenolato)aluminum]bis $[\mu$ -(3,5-bis(trifluoromethyl)-thiophenolato)]bis(triisobutylphosphine)-copper;

[μ-(3,5-bis(trifluoromethyl)-thiophenolato)]bis(trihexylphosphine)-copper;

[bis(3,5-bis(trifluoromethyl)-thiophenolato)aluminum]bis [μ-(3,5-bis(trifluoromethyl)-thiophenolato)]bis(triphenylphosphine)-copper;

[bis(3,5-bis(trifluoromethyl)-thiophenolato)aluminum]bis [μ-(3,5-bis(trifluoromethyl)-thiophenolato)]bis(perfluorotriphenylphosphine)-copper;

[bis(thiophenolato)aluminum]bis[\u03c4-(thiophenolato)]bis(triisobutylphosphine)-copper;

[bis(thiophenolato)aluminum]bis[μ-(thiophenolato)]bis(trihexylphosphine)-copper;

 $[bis(thiophenolato)aluminum]bis[\mu-(thiophenolato)]bis$ (triphenylphosphine)-copper;

35 [bis(thiophenolato)aluminum]bis[μ-(thiophenolato)]bis(perfluorotriphenylphosphine)-copper;

[bis(α -toluenethiolato)aluminum]bis[μ -(α -toluenethiolato)] bis(triisobutylphosphine)-copper;

[bis(α -toluenethiolato)aluminum]bis[μ -(α -toluenethiolato)] bis(trihexylphosphine)-copper;

[bis(α -toluenethiolato)aluminum]bis[μ -(α -toluenethiolato)] bis(triphenylphosphine)-copper; [bis(α -toluenethiolato) aluminum]bis[μ -(α -toluenethiolato)]bis(perfluorotriphenylphosphine)-copper;

45 [bis(pentafluorothiophenolato)aluminum]bis[μ-(pentafluorothiophenolato)]bis(triisobutylphosphine)-copper; [bis (pentafluorothiophenolato)aluminum]bis[u-(pentafluorothiophenolato)]bis(trihexylphosphine)-copper;

[bis(pentafluorothiophenolato)aluminum]bis[μ-(pentafluorothiophenolato)]bis(triphenylphosphine)-copper; [bis(pentafluorothiophenolato)aluminum]bis[µ-(pentafluorothiophenolato)]bis(perfluorotriphenylphosphine)-

By way of example and not limitation, the following are 55 examples of silver-aluminum SSPs:

[bis(ethanethiolato)aluminum]bis[µ-(ethanethiolato)]bis(triisobutylphosphine)-silver;

 $[bis(ethanethiolato) a luminum] bis [\mu-(ethanethiolato)] bis(tri-luminum) bis [\mu-(ethanethiolato)] bis(tri-luminum) bis[\mu-(ethanethiolato)] bis(tri-luminum) bis(\mu-(ethanethiolato)) bis(tri-luminum) bis(\mu-(ethanethiolato)) bis(tri-luminum) bis(\mu-(ethanethiolato)) b$ hexylphosphine)-silver;

[bis(ethanethiolato)aluminum]bis[µ-(ethanethiolato)]bis (triphenylphosphine)-silver;

[bis(ethanethiolato)aluminum]bis[μ-(ethanethiolato)]bis (perfluorotriphenylphosphine)-silver;

 $[bis(propanethiolato)aluminum]bis[\mu-(propanethiolato)]bis$ (triisobutylphosphine)-silver;

[bis(propanethiolato)aluminum]bis[μ-(propanethiolato)]bis (trihexylphosphine)-silver;

[bis(propanethiolato)aluminum]bis[μ-(propanethiolato)]bis (triphenylphosphine)-silver;

[bis(propanethiolato)aluminum]bis[μ-(propanethiolato)]bis (perfluorotriphenylphosphine)-silver;

[bis(hexanethiolato)aluminum]bis[μ-(propanethiolato)]bis (triisobutylphosphine)-silver;

[bis(hexanethiolato)aluminum]bis[μ-(propanethiolato)]bis (trihexylphosphine)-silver;

[bis(hexanethiolato)aluminum]bis[μ-(propanethiolato)]bis (triphenylphosphine)-silver;

[bis(hexanethiolato)aluminum]bis[µ-(propanethiolato)]bis (perfluorotriphenylphosphine)-silver;

[bis(4-trifluoromethyl-thiophenolato)aluminum]bis[\mu-(4-trifluoromethylthiophenolato)]bis(triisobutylphosphine)-silver:

 $[bis (4-trifluoromethyl-thiophenolato) a luminum] bis [\mu-(4-trifluoromethyl-thiophenolato)] bis (trihexylphosphine)-silver:$

[bis(4-trifluoromethyl-thiophenolato)aluminum]bis[µ-(4-trifluoromethyl-thiophenolato)]bis(triphenylphosphine)-silver:

[bis(4-trifluoromethyl-thiophenolato)aluminum]bis[μ-(4-trifluoromethyl-thiophenolato)]bis(perfluorotriphenylphosphine)-silver;

[bis(3,5-bis(trifluoromethyl)-thiophenolato)aluminum]bis [μ -(3,5-bis(trifluoromethyl)-thiophenolato)]bis(triisobutylphosphine)-silver;

[bis(3,5-bis(trifluoromethyl)-thiophenolato)aluminum]bis $[\mu$ -(3,5-bis(trifluoromethyl)-thiophenolato)]bis(trihexylphosphine)-silver;

 $[bis (3,5-bis (trifluoromethyl)-thiophenolato) aluminum] bis \\ [\mu-(3,5-bis (trifluoromethyl)-thiophenolato)] bis (triphenylphosphine)-silver;$

[bis(3,5-bis(trifluoromethyl)-thiophenolato)aluminum]bis [μ-(3,5-bis(trifluoromethyl)-thiophenolato)]bis(perfluorotriphenylphosphine)-silver;

[bis(thiophenolato)aluminum]bis[μ-(thiophenolato)]bis(triisobutylphosphine)-silver;

 $[bis (thiophenolato) a luminum] bis [\mu-(thiophenolato)] bis (trihexylphosphine)-silver;$

[bis(thiophenolato)aluminum]bis[μ-(thiophenolato)]bis (triphenylphosphine)-silver;

[bis(thiophenolato)aluminum]bis[μ -(thiophenolato)]bis(perfluorotriphenylphosphine)-silver; [bis(α -toluenethiolato) 45 aluminum]bis[μ -(α -toluenethiolato)]bis(triisobutylphosphine)-silver; [bis(α -toluenethiolato)aluminum]bis[μ -(α -toluenethiolato)]bis(trihexylphosphine)-silver; [bis(α -toluenethiolato)]bis (triphenylphosphine)-silver; [bis(α -toluenethiolato)]bis (triphenylphosphine)-silver; [bis(α -toluenethiolato)]bis (perfluorotriphenylphosphine)-silver;

[bis(pentafluorothiophenolato)aluminum]bis[μ-(pentafluorothiophenolato)]bis(triisobutylphosphine)-silver; [bis (pentafluorothiophenolato)aluminum]bis[μ-(pentafluorothiophenolato)]bis(trihexylphosphine)-silver;

[bis(pentafluorothiophenolato)aluminum]bis[μ-(pentafluorothiophenolato)]bis(triphenylphosphine)-silver; and [bis (pentafluorothiophenolato)aluminum]bis[μ-(pentafluorothiophenolato)]bis(perfluorotriphenylphosphine)-silver. The examples of SSPs set forth above are examples only, and those of ordinary skill in the art will understand that such examples may be used to derive many other SSPs for use in forming chalcogenide materials such as, for example, ternary chalcopyrite materials.

Embodiments of the present invention include methods of forming SSPs. The methods are believed to be generally 12

faster, more versatile (may be used to form a wider variety of SSPs), and to exhibit improved yield relative to other methods known in the art.

An example of a reaction pathway in accordance with some embodiments of the present invention is represented by Reactions 1 through 3 below:

In some embodiments of the present invention, SSPs of the general formula $L_2N(\mu\text{-ER})_2M'(\text{ER})_2$ may be formed using a reaction pathway in accordance with Reactions 1 through 3 below:

$$N-X+2L\rightarrow L_2N-X$$
 Reaction 1

$$L_2N-X+M'X_3 \rightarrow \frac{1}{2}\{L_2N(\mu-X)_2M'X_2\}_2$$
 Reaction 2

$$\frac{1}{2}$$
{L₂N(μ-X)₂M'X₂}₂+4MER \rightarrow L₂N(μ-ER)₂M'(ER)₂
(+4MX)(SSP)
Reaction 3

wherein L is a Lewis base (that is coordinated to N by a dative bond), each N is individually selected from Group IB atoms, 20 each M is individually selected from Group IA atoms, each M' is individually selected from Group IIIA atoms, each E is individually selected from Group VIA atoms, each X is individually selected from Group VIIA atoms or a nitrate group, and each R group is individually selected from the group consisting of alkyl, aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl, silane, and carbamato groups. In accordance with some embodiments of the present invention, the Lewis base L may comprise, for example, a substituted phosphine of the general formula PR₃, wherein P is a phosphorous atom and each R is individually selected from the group consisting of alkyl, aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl, silane, and carbamato groups. As a non-limiting example, the Lewis base L may comprise a phosphine compound such as Ph₃P, wherein Ph is the phenyl group (a functional group comprising an 35 aromatic cyclic ring of the formula C₆H₅⁻). In accordance with additional embodiments of the present invention, the Lewis base L may comprise, for example, a substituted multidentate phosphine of the general formula R₂PAPR₂, wherein P is a phosphorous atom, each A is individually selected from the group consisting of alkyl, aryl, and vinyl, and each R is individually selected from the group consisting of alkyl, aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl, silane, and carbamato groups. As a non-limiting example, the Lewis base L may comprise a phosphine compound such as Ph₂PC₆H₄OC₆H₄PPh₂ (bis(2-diphenylphosphinophenyl) ether), wherein Ph is the phenyl group (a functional group comprising an aromatic cyclic ring of the formula C₆H₅⁻).

Reactions 1 through 3 above, when combined and balanced, result in the net reaction $N-X+2L+M'X_3+4MER\rightarrow L_2N(\mu-ER)_2M'(ER)_2(+4MX)$. The particular reaction pathway prescribed by Reactions 1 through 3 results in the formation of the intermediate products L_2N-X and $\frac{1}{2}\{L_2N(\mu-X)_2M'X_2\}_2$. Such reaction pathways provide certain advantages over previously known methods of forming SSPs having the general formula $L_2N(\mu-ER)_2M'(ER)_2$.

Referring to Reaction 1, a first reagent comprising L_2N —X (for use in Reaction 2) may be formed by reacting a Lewis base (e.g., a substituted phosphine (PR $_3$)) with a metal halide or nitride (N—X). The molar ratio of the Lewis base to the metal halide or nitride may be, for example, two (2) to one (1) (in other words, two molar equivalents of the Lewis base and one molar equivalent of the metal halide or nitride may be reacted with one another in Reaction 1), although the particular molar ratio may depend on the particular SSP to be formed.

Reaction 1 may be carried out in solution. For example, Reaction 1 may be performed in anhydrous benzene (C₆H₆),

in acetonitrile (CH₃CN), in a mixture of more than one solvent, or in another suitable solvent. The product of Reaction 1 may comprise a suspension in which the first reagent comprising L₂N—X is present as a liquid or solid product suspended in the solution. The suspension may be concentrated 5 to yield a liquid or a solid product comprising the L₂N—X substance. Alternatively, the reaction may be conducted in a single pot reaction, wherein the L₂N—X substance is not concentrated.

Referring to Reaction 2, the intermediate product ${}^{1}\!\!/_2\{L_2N \ 10 \ (\mu\text{-}X)_2\text{M}'X_2\}_2}$ may be formed by reacting the first reagent comprising L_2N —X with $M'X_3$. The molar ratio of the L_2N —X reagent to the $M'X_3$ reagent may be, for example, one (1) to one (1) (in other words, one molar equivalent of the L_2N —X reagent and one molar equivalent of the $M'X_3$ 15 reagent may be reacted with one another in Reaction 2), although, again, the particular molar ratio may depend on the particular SSP to be formed.

Reaction 2 also may be carried out in solution. For example, Reaction 2 may be performed in the same benzene 20 (C_6H_6) (or other solvent or solvent mixture) used in Reaction 1 or another suitable solvent (i.e., a solvent in which the L_2N-X reagent and/or the M'X₃ reagent is soluble). The product of Reaction 2 may comprise a solution or a suspension comprising the intermediate product $\frac{1}{2}\{L_2N(\mu-X)_2\}$ 25 $\frac{1}{2}$

Referring to Reaction 3, an SSP of the general formula $L_2N(\mu-ER)_2M'(ER)_2$ may be formed by reacting the intermediate product $\frac{1}{2}\{L_2N(\mu-X)_2M'X_2\}_2$ of Reaction 2 with the MER reagent. By way of example and not limitation, the 30 MER reagent may be added to the solution or suspension that is the product of Reaction 2 and includes the intermediate product $\frac{1}{2}\{L_2N(\mu-X)_2M'X_2\}_2$. The molar ratio of the MER reagent to the $\frac{1}{2}\{L_2N(\mu-X)_2M'X_2\}_2$ reagent may be, for example, four (4) to one (1) (in other words, four molar 35 equivalents of the MER reagent and one molar equivalent of the $\frac{1}{2}\{L_2N(\mu-X)_2M'X_2\}_2$ reagent may be reacted with one another in Reaction 3), although, again, the particular molar ratio may depend on the particular SSP to be formed. It is noted that the R groups in the four molar equivalents of the 40 MER reagent may differ from one another, and may be individually selected from the group consisting of alkyl, aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl, silane, and carbamato groups. In other words, one molar equivalent of the MER reagent may comprise R¹, one molar equivalent of the 45 MER reagent may comprise R², one molar equivalent of the MER reagent may comprise R³, and one molar equivalent of the MER reagent may comprise R⁴, wherein R¹, R², R³, and R⁴ all differ from one another. In additional embodiments, less than four different R groups (e.g., one, two, or three), or 50 more than four different R groups may be employed in the MER reagent. Similarly, the E groups in the four molar equivalents of the MER reagent may differ from one another, and may be individually selected from the Group VIA atoms. In other words, one molar equivalent of the MER reagent may 55 comprise E1, one molar equivalent of the MER reagent may comprise E², one molar equivalent of the MER reagent may comprise E³, and one molar equivalent of the MER reagent may comprise E^4 , wherein E^1 , E^2 , E^3 , and E^4 all differ from one another. In additional embodiments, less than four differ- 60 ent E groups (e.g., one, two, or three), or more than four different E groups may be employed in the MER reagent.

Reaction 3 also may be carried out in solution. For example, Reaction 3 may be performed in the same in the same benzene (C₆H₆) (or other solvent or solvent mixture) 65 used in Reaction 2 or another suitable solvent, after carrying out Reaction 2, such that the solution includes the intermedi-

14

ate product and reagent $\frac{1}{2}\{L_2N(\mu-X)_2M'X_2\}_2$. The product of Reaction 3 may comprise a suspension in which the SSP of the general formula $L_2N(\mu-ER)_2M'(ER)_2$ is present as a liquid or solid product dissolved or suspended in the solution. The by-product may be filtered, and the filtrate may be concentrated to isolate the SSP. The SSP may comprise a liquid or solid product, and the solid SSP may be re-crystallized as desirable in preparation for utilizing the SSP to form a chalcogenide material (e.g., a ternary chalcopyrite material).

In accordance with some embodiments of the present invention, SSPs of the general formula $L_2N(\mu\text{-}E^1R^1)(\mu\text{-}E^2R^2)$ $M'(E^2R^2)_2(SSP)$ or $L_2N(\mu\text{-}E^2R^2)_2M'(E^2R^2)(E^1R^1)$ (SSP) may be formed using reaction pathways in accordance with additional embodiments of the present invention represented by Reaction 1 (which is the same as above and repeated below for convenience) and Reactions 4 through 6 below:

N—X+2L
$$\to$$
L₂N—X Reaction 1
L₂N—X+ME¹R¹ \to L₂N-E¹R¹(+MX) Reaction 4
M'X₃+nME²R² \to M'(E²R²)_n(+nMX) Reaction 5
L₂N-E¹R¹+M'(E²R²)_n \to L₂N(μ -E¹R¹)(μ -E²R²)M' (E²R²)₂(SSP) or L₂N(μ -E²R²)₂M'(E²R²)(E¹R¹) (SSP)

wherein L is a Lewis base (that is coordinated to N by a dative bond), each N is individually selected from Group IB atoms, each M is individually selected from Group IIA atoms, each M' is individually selected from Group IIIA atoms, each E¹ and E² is individually selected from Group VIA atoms (E¹ and E² may be same or different), each X is individually selected from Group VIIA atoms or a nitrate group, n is any number (e.g., an integer) from one (1) to three (3), and each R¹ and R² group is individually selected from the group consisting of alkyl, aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl, silane, and carbamato groups, wherein at least one E¹ differs from at least one E² and/or at least one R¹ differs from at least one R². In some embodiments, each R¹ may differ from each R².

Reaction 1 may be carried out as previously described herein. Referring to Reaction 4, the intermediate product L_2N - E^1R^1 may be formed by reacting the L_2N -X (obtained by Reaction 1) with ME^1R^1 . The molar ratio of the L_2N -X reagent to the ME^1R^1 reagent may be, for example, one (1) to one (1) (in other words, one molar equivalent of the L_2N -X reagent and one molar equivalent of the ME^1R^1 reagent may be reacted with one another in Reaction 4), although, again, the particular molar ratio may depend on the particular SSP to be formed.

Reaction 4 also may be carried out in solution. For example, Reaction 4 may be performed in the same benzene (C_6H_6) or other solution used in Reaction 1, or in another suitable solvent (i.e., a solvent in which the L_2N —X reagent and/or the ME^1R^1 reagent is soluble). The product of Reaction 4 may comprise a solution or a suspension comprising the intermediate product L_2N - E^1R^1 .

Referring to Reaction 5, the intermediate product $M'(E^2R^2)_m$ may be formed by reacting $M'X_3$ with ME^2R^2 . The molar ratio of the $M'X_3$ reagent to the ME^2R^2 reagent may be, for example, one (1) to three (3) (in other words, one molar equivalent of the $M'X_3$ reagent and three molar equivalents of the ME^2R^2 reagent may be reacted with one another in Reaction 5), although, again, the particular molar ratio may depend on the particular SSP to be formed.

Reaction 5 also may be carried out in solution. For example, Reaction 5 may be performed in the same benzene (C_6H_6) or other solution (i.e., a solvent in which the M'X₃

reagent and/or the ME^2R^2 reagent is soluble). The product of Reaction 5 may comprise a solution or a suspension comprising the intermediate products $M'(E^2R^2)_n$. It is noted that the R^2 groups in the four molar equivalents of the ME^2R^2 reagent may be individually selected from the group consisting of 5 alkyl, aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl, silane, and carbamato groups. Thus, each of the R^2 groups may be the same or they may differ from one another. Similarly, the E^2 groups in the four molar equivalents of the ME^2R^2 reagent may be individually selected from the Group V1A atoms, and, 10 thus, may be the same or they may differ from one another.

Referring to Reaction 6, an SSP of the general formula $L_2N(\mu\text{-}E^1R^1)(\mu\text{-}E^2R^2)M'(E^2R^2)_2$ or $L_2N(\mu\text{-}E^2R^2)_2M'(E^1R^1)$ (E^2R^2) may be formed by reacting the intermediate product $L_2N\text{-}E^1R^1$ of Reaction 4 with the intermediate product $M'(E^2R^2)_n$ of Reaction 5. By way of example and not limitation, the intermediate product $M'(E^2R^2)_n$ may be added to the solution or suspension that is the product of Reaction 4 and includes the intermediate product $L_2N\text{-}E^1R^1$. The molar ratio of the $L_2N\text{-}E^1R^1$ reagent to the $M'(E^2R^2)_n$ reagent may be, for example, one (1) to one (1) (in other words, one molar equivalent of the $L_2N\text{-}E^1R^1$ reagent and one molar equivalent of the $M'(E^2R^2)_n$ reagent may be reacted with one another in Reaction 6), although, again, the particular molar ratio may depend on the particular SSP to be formed.

Reaction 6 also may be carried out in solution. For example, Reaction 6 may be performed in the same benzene (C_6H_6) or other solution used in Reaction 5, after carrying out Reaction 5, such that the solution includes the intermediate product and reagent $M'(E^2R^2)$. The product of Reaction 6 30 may comprise a suspension in which the SSP of the general formula $L_2N(\mu\text{-}E^1R^1)(\mu\text{-}E^2R^2)M'(E^2R^2)_2$ or $L_2N(\mu\text{-}E^2R^2)_2$ $M'(E^1R^1)(E^2R^2)$ is present as a liquid or solid dissolved or suspended in the solution. The by-product may be filtered, and the filtrate may be concentrated to isolate the SSP. The 35 SSP may comprise a liquid or solid product, and the solid SSP may be re-crystallized as desirable in preparation for utilizing the SSP to form a chalcogenide material (e.g., a ternary chalcopyrite material).

In accordance some embodiments of the present invention, 40 SSPs of the general formula $L_2N(\mu-ER^1)(\mu-ER^2)M'(ER^3)$ (ER⁴) may be formed using Reactions 1 and 4 (which are the same as above and repeated below for convenience) and Reactions 7 through 10 below:

$$\begin{array}{ll} L_2N(\mu\text{-ER}^1)(\mu\text{-X})M'X_2+MER^2L_2N(\mu\text{-ER}^1)(\mu\text{-ER}^2) \\ M'X_2(+MX) \end{array}$$
 Reaction 8

$$\begin{array}{ll} L_2N(\mu\text{-ER}^1)(\mu\text{-ER}^2)M'X_2+MER^3 \!\rightarrow\! L_2N(\mu\text{-ER}^1)(\mu\text{-}\\ ER^2)M'(ER^3)X(+MX) \end{array}$$
 Reaction 9 55

$$\begin{array}{ll} L_2N(\mu\text{-ER}^1)(\mu\text{-ER}^2)M'(ER^3)X + MER^4 {\to} L_2N(\mu\text{-ER}^1) \\ (\mu\text{-ER}^2)M'(ER^3)(ER^4)(+MX)(SSP) \end{array} \qquad \qquad \text{Reaction } 10$$

wherein L is a Lewis base that is coordinated to N by a dative bond, each N is individually selected from Group 60 1B atoms, each M is individually selected from Group 1A atoms, each M' is individually selected from Group 111A atoms, each E is individually selected from Group V1A atoms, each X is individually selected from Group V11A atoms or a nitrate group, each R¹, R², R³, and R⁴ group 65 is individually selected from the group consisting of alkyl, aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl,

16

silane, and carbamato groups, and R^1 , R^2 , R^3 , R^4 are different from one another. The E groups in the MER¹, MER², MER³, and MER⁴ reagents may be the same or they may be different from one another. In other words, the MER¹ reagent may comprise E^1 , the MER² reagent may comprise E^3 , and the MER⁴ reagent may comprise E^4 , wherein E^1 , E^2 , E^3 , and E^4 all differ from one another. In additional embodiments, less than four different E groups (e.g., one, two, or three), or more than four different E groups may be employed in the MER¹, MER², MER³, and MER⁴ reagents.

Reactions 1 and 4 may be carried out as previously described herein.

Referring to Reaction 7, the intermediate product $L_2N(\mu-ER^1)(\mu-X)M'X_2$ (which may be an SSP) may be formed by reacting the L_2N-ER^1 (obtained by Reaction 4) with $M'X_3$. The molar ratio of the L_2N-ER^1 reagent to the $M'X_3$ reagent may be, for example, one (1) to one (1) (in other words, one molar equivalent of the $M'X_3$ reagent and one molar equivalent of the L_2N-ER^1 reagent may be reacted with one another in Reaction 7). Reaction 7 may be carried out in a solution such as benzene (C_6H_6) or in another suitable solvent or mixture of solvents. The product of Reaction 7 may comprise a solution or a suspension comprising the intermediate SSP product $L_2N(\mu-ER^1)(\mu-X)M'X_2$.

Referring to Reaction 8, the intermediate SSP product $L_2N(\mu-ER^1)(\mu-ER^2)M'X_2$ may be formed by reacting one (1) molar equivalent of MER² with the intermediate SSP product $L_2N(\mu-ER^1)(\mu-X)M'X_2$ of Reaction 7. As shown in Reaction 9, the intermediate SSP product L₂N(μ-ER¹)(μ-ER²)M' (ER³)X may be formed by reacting one (1) molar equivalent of MER³ with one (1) molar equivalent of the SSP intermediate product $L_2N(\mu-ER^2)(\mu-ER^2)M'X_2$ obtained by Reaction 8. Similarly, as shown in Reaction 10, a final SSP product of the general formula $L_2N(\mu-ER^1)(\mu-ER^2)M'(ER^3)(ER^4)$ may be formed by reacting one (1) molar equivalent of MER⁴ with one (1) molar equivalent of the SSP intermediate product $L_2N(\mu-ER^1)(\mu-ER^2)M'(ER^3)X$ obtained by Reaction 9. The product of Reaction 10 may comprise a suspension in which an SSP of the general formula $L_2N(\mu\text{-ER}^1)(\mu\text{-ER}^2)M'(ER^3)$ (ER⁴) is present as a liquid or solid dissolved or suspended in the solution. It is noted that the various groups may not be added to the particular positions of the SSP in the order represented above, and, thus, the ER1, ER2, ER3, and ER4 groups may be in any position in the SSP. The by-product may be filtered, and the filtrate may be concentrated to isolate the SSP. The SSP may comprise a liquid or solid product, and the solid SSP may be re-crystallized as desirable in preparation 50 for utilizing the SSP to form a chalcogenide material (e.g., a ternary chalcopyrite material).

In accordance with additional embodiments of the present invention, after forming SSPs as previously described herein, the composition of the SSPs may be tailored by carrying out one or more additional reactions (e.g., substitution reactions) with the SSPs, as described below. For example, after forming a first SSP (SSP¹), a second, different SSP (SSP²) of the general formula $L_2N(\mu\text{-}E^1R^1)(\mu\text{-}E^2R^2)M'(E^3R^3)(E^4R^4)$ may be formed from the first SSP¹ in accordance with Reaction 11 below:

$$SSP^1 + nHE^1R^1 + mHE^2R^2 + oHE^3R^3 + pHE^4R^4 \rightarrow L_2N(\mu-E^1R^1)(\mu-E^2R^2)M'(E^3R^3)(E^4R^4)(SSP^2) \\ \qquad \qquad Reaction \ 11$$

wherein each of n, m, o, and p is any number between zero (0) and four (4), the sum of n, m, o, and p is four (4), L is a Lewis base that is coordinated to N by a dative bond, each N is individually selected from Group IB atoms, each M' is indi-

vidually selected from Group 111A atoms, each of E^1, E^2, E^3 , and E^4 is individually selected from Group V1A atoms, each of R^1, R^2, R^3 , and R^4 is individually selected from the group consisting of alkyl, aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl, silane, and carbamato groups. In Reaction 11, R^1, R^2, S^3 , and R^4 may be the same or different from one another, and E^1, E^2, E^3 , and E^4 may be the same or different from one another.

As one non-limiting example, after forming an SSP of the general formula $L_2N(\mu-ER)_2M'(ER)_2$ using Reactions 1 10 through 3, as previously described, a different SSP of the general formula $L_2N(\mu-E^1R^1)_2M'(E^1R^1)_2$ may be formed from the SSP in accordance with Reaction 11 by reacting four (4) molar equivalents of HE¹R¹ with one (1) molar equivalent of the $L_2N(\mu-E^1R^1)_2M'(E^1R^1)_2SSP$. As another non-limiting 1 example, after forming an SSP of the general formula L₂N $(\mu$ -E¹R¹) $(\mu$ -E²R²)M'(E²R²)₂ using Reactions 1, 4, 5, and 6, as previously described, a different SSP of the general formula $L_2N(\mu-E^3R^3)_2M'(E^3R^3)_2$ may be formed from the SSP in accordance with Reaction 11 by reacting four (4) molar 20 equivalents of HE³R³ with one (1) molar equivalent of the $L_2N(\mu-E^1R^1)(\mu-E^2R^2)M'(E^2R^2)_2SSP$. As yet another nonlimiting example, after forming an SSP of the general formula $L_2N(\mu-ER^1)(\mu-ER^2)M'(ER^3)(ER^4)$ using Reactions 1, 4, and 7 through 10, as previously described, a different SSP of the 25 general formula $L_2N(\mu\text{-}E^2R^5)_2M'(E^2R^5)_2$ may be formed from the SSP in accordance with Reaction 11 by reacting four (4) molar equivalents of HE²R⁵ with one (1) molar equivalent of the $L_2N(\mu-ER^1)(\mu-ER^2)M'(ER^3)(ER^4)SSP$.

Reaction 11 also may be carried out in solution. For 30 example, Reaction 11 may be performed in benzene (C_6H_6) or another suitable solvent or solvent mixture. The product of Reaction 11 may comprise a suspension in which the SSP product is present as liquid or solid matter dissolved or suspended in the solution. The by-product may be filtered, and 35 the filtrate may be concentrated to isolate the SSP product. The SSP product may comprise a liquid or solid product, and the solid SSP may be re-crystallized as desirable in preparation for utilizing the SSP to form a chalcogenide material (e.g., a ternary chalcopyrite material).

The reaction mechanisms described hereinabove with reference to Reactions 1 through 11 may be used to form a wide variety of SSPs. For example, different species of the L₂N—X reagent (Reaction 2) and the ME¹R¹ reagent (Reaction 3) may be used to place different R¹ groups on the SSPs, 45 different E¹ groups on the SSPs, or different L groups on the SSPs, and different species of the L₂N—X reagent, the M'X₃ reagent, and the ME¹R¹ reagent may be used in selected ratios to tailor the identity and concentration of the N, M', and E atoms in the SSPs. Four non-limiting examples of embodi- 50 ments of the present invention are set forth below to illustrate the versatility in forming SSPs in accordance with embodiments of the present invention. Embodiments of the present invention, however, are not to be limited to methods of forming the specific SSPs formed in the examples below, or to 55 methods employing the particular reagents employed in the examples below.

Example 1

The SSP $(Ph_3P)_2Cu(\mu\text{-SEt})_2ln(SEt)_2$ may be formed as follows.

Two (2) molar equivalents of Ph_3P (42.000 grams, 160.13 millimoles) may be added to a mixture of two hundred forty (240) milliliters (mL) of anhydrous benzene (C_6H_6) and two 65 hundred forty (240) milliliters (mL) of anhydrous tetrahydrofuran ((CH_2)₄O, THF) to form a first solution, which may be

18

stirred. One (1) molar equivalent of anhydrous Cu(1)Cl (7.9260 grams, 80.064 millimoles) may be added to the stirring first solution to form a first white suspension including the intermediate product (Ph_3P)₂Cu—Cl. The first suspension may be concentrated to form a solid product or kept to continue the reaction as a one pot reaction, which is referred to herein as "Mixture A."

A second solution may be formed by adding 1.4770 molar equivalents of KH (40.000 grams, 299.18 millimoles, collected from thirty (30) weight percent dispersion in mineral oil and washed several times with benzene) to about 100 mL of benzene. One (1) molar equivalent of EtSH (15.000 mL, 202.56 millimoles) (Et is the ethyl group) may be added to the second solution and stirred for a few minutes. The EtSH and the KH react with one another to form KSEt. The resulting product that includes the KSEt (which may comprise a solution or a solid product comprising the KSEt formed by drying a solution) is referred to herein as "Mixture B."

A "Mixture C" may also be formed by adding one (1) molar equivalent of Na metal (34.000 grams, 1478.9 millimoles, collected from mineral oil and washed several times with diethyl ether) to about 500 mL of diethyl ether. One (1) molar equivalent of EtSH (109.52 mL, 1478.9 millimoles) may be added to the second solution and stirred for twelve (12) hours at forty (40) degrees Celsius (° C.). The EtSH and the Na metal react with one another to form NaSEt. The resulting product that includes the NaSEt (which may comprise a solution or a solid product comprising the NaSEt formed by drying a solution) is Mixture C.

If the reaction is a one pot reaction, one (1) molar equivalent of anhydrous $InCl_3$ (17.709 grams, 80.064 millimoles) may be added to Mixture A, after which this third solution may be stirred for about sixty (60) minutes at eighty (80) degrees Celsius (° C.). The resulting mixture may comprise a fourth solution. This solution may comprise the intermediate product $\{(Ph_3P)_2Cu(\mu-Cl)_2InCl_2\}_2$, a computer generated graphical representation of which is illustrated in FIG. 1. As shown in FIG. 1, this intermediate product comprises an eight (8) member ring structure defined by two (2) copper atoms, two (2) indium atoms, and four (4) chlorine atoms.

Four (4) molar equivalents of NaSEt (26.940 grams, 320.26 millimoles) or KSEt (32.100 grams, 320.26 millimoles) provided by Mixture B or C, respectively, may be added to the fourth solution and stirred for an additional twelve (12) hours at eighty (80)° C.

In some embodiments, however, less than four (4) molar equivalents of the NaSEt or KSEt may initially be added to the fourth solution. Fractions of the four (4) molar equivalents may be added sequentially to attain various intermediate products. For example, one (1) molar equivalent of the NaSEt or KSEt may initially be added to the fourth solution to form an intermediate product, the exact nature of which is not known, but is currently believed to be the (Ph₃P)₂Cu(μ-SEt) lnCl₃ species shown in the computer generated graphical representation of F1G. 2. A second molar equivalent of the NaSEt or KSEt may be added to the fourth solution to form another intermediate product, the exact nature of which is not known, but is currently believed to be the (Ph₃P)₂Cu(μ-SEt) 2lnCl₂ species shown in the computer generated graphical representation of F1G. 3. Similarly, a third molar equivalent of the NaSEt or KSEt may then be added to the fourth solution to form another intermediate product, the exact nature of which is not known, but is currently believed to be the (Ph₃P) ₂Cu(μ-SEt)₂In(SEt)Cl species shown in the computer generated graphical representation of FIG. 4. Finally, a fourth molar equivalent of the NaSEt may be added to the fourth

solution to form the final SSP product $(Ph_3P)_2Cu(\mu\text{-SEt})_2ln$ (SEt)₂, which is shown in the computer generated graphical representation of FIG. **5**.

After all four (4) molar equivalents of the NaSEt or KSEt provided by Mixture B or C, respectively, have been added to the fourth solution, the solution may be stirred for an additional twelve (12) hours at eighty degrees Celsius (80° C.). After stirring, the resulting mixture may comprise a suspension that includes a solution with particles of the SSP (Ph₃P)₂ Cu(μ-SEt)₂ln(SEt)₂ suspended therein. The SSP may be isolated by filtering of by-product, and the filtrate may be concentrated as necessary or desirable. Optionally, the isolated SSP may be re-crystallized in preparation for using the SSP to form the ternary chalcopyrite material CulnS₂.

It will be apparent from the above description that, in additional embodiments, instead of using the same reagent (e.g., NaSEt) in each of the sequential additions to the reaction mixture, different reagents including different moieties (other than ethyl groups) may be added to the fourth solution 20 to substitute different, selected moieties into the resulting SSP product at different selected locations within the structure of the SSP molecule or complex.

Example 2

The SSP $(Ph_3P)_2Cu(\mu\text{-SBn})(\mu\text{-SEt})Ga(SEt)_2$ may be formed as follows.

The intermediate product $(Ph_3P)_2Cu$ —Cl is formed as described in Example 1 above (and repeated below for convenience). Two (2) molar equivalents of Ph_3P (42.000 grams, 160.13 millimoles) may be added to mixture of two hundred forty (240) milliliters (mL) of anhydrous benzene (C_6H_6) and two hundred forty (240) milliliters (mL) of anhydrous tetrahydrofuran (($CH_2)_4O$, THF) to form a first solution, which 35 may be stirred. One (1) molar equivalent of anhydrous Cu(1) Cl (7.9260 grams, 80.064 millimoles) may be added to the stirring first solution to form a first white suspension including the intermediate product $(Ph_3P)_2Cu$ —Cl. The first suspension may be concentrated to form a solid product or kept 40 to continue the reaction, which is referred to herein as "Mixture A."

A second solution (which is the same as above and repeated below for convenience) may be formed by adding one (1) molar equivalent of Na metal (34.000 grams, 1478.9 milli-45 moles, collected from mineral oil and washed several times with diethyl ether) to about 500 mL of diethyl ether. One (1) molar equivalent of EtSH (109.52 mL, 1478.9 millimoles) may be added to the second solution and stirred for twelve (12) hours at forty (40) degrees Celsius (° C.). The EtSH and 50 the Na metal react with one another to form NaSEt. The resulting product that includes the NaSEt (which may comprise a solution or a solid product comprising the NaSEt formed by drying a solution) is referred to herein as "Mixture B."

A third solution may be formed by adding one (1) molar equivalent of Na metal (10.000 grams, 434.97 millimoles, collected from mineral oil and washed several times with diethyl ether) to about 200 mL of diethyl ether. One (1) molar equivalent of $C_6H_5CH_2SH$ (HSBn, 51.066 mL, 434.97 millimoles) may be added to the third solution and stirred for twelve (12) hours at 40° C. The $C_6H_5CH_2SH$ and the Na metal react with one another to form NaSCH $_2C_6H_5$ (NaSBn). The resulting product that includes the NaSBn (which may comprise a solution or a solid product comprising the NaSBn 65 formed by drying a solution) is referred to herein as "Mixture C"

20

A fourth solution may be formed by adding one (1) molar equivalent of $(Ph_3P)_2Cu$ —Cl (49.926 grams, 80.064 millimoles) provided by Mixture A to about 200 mL of benzene. One (1) molar equivalent of NaSBn (8.1840 grams, 80.064 millimoles) provided by Mixture C may be added to the fourth solution and stirred for twelve (12) hours. The $(Ph_3P)_2$ Cu—Cl and the NaSBn react with one another to form $(Ph_3P)_2$ CuSBn. The resulting mixture may comprise a fourth solution. This solution may comprise the intermediate product $(Ph_3P)_2$ CuSBn (which may comprise a solution or a solid product comprising the $(Ph_3P)_2$ CuSBn formed by drying a solution) is referred to herein as "Mixture D."

A fifth solution may be formed by adding three (3) molar equivalents of NaSEt (20.205 grams, 240.192 millimoles) provided by Mixture B to about 200 mL of benzene. One (1) molar equivalent of anhydrous GaCl₃ (14.098 grams, 80.064 millimoles) may be added to this fifth solution, after which the fifth solution may be stirred for about sixty (60) minutes at eighty (80)° C. The resulting mixture may comprise a fifth solution. This solution may comprise the intermediate product Ga(SEt)₃ (which may comprise a solution or a solid product comprising the Ga(SEt)₃ formed by drying a solution).

One (1) molar equivalent of (Ph₃P)₂CuSBn (56.950 grams, 80.064 millimoles) provided by Mixture D may be added to 25 the fifth solution and stirred for an additional twelve (12) hours at eighty (80)° C. After stirring, the resulting mixture may comprise a suspension that includes a solution with particles of the SSP (Ph₃P)₂Cu(μ-SBn)(μ-SEt)Ga(SEt)₂ suspended therein. The SSP may be isolated by filtering of by30 product, and the filtrate may be concentrated as necessary or desirable. Optionally, the isolated SSP may be re-crystallized in preparation for using the SSP to form the ternary chalcopyrite material CuGaS₂.

In some embodiments, however, less than three (3) molar equivalents of the NaSEt may initially be added to the fifth solution. Fractions of the three (3) molar equivalents may be added sequentially to attain various intermediate products. For example, one (1) molar equivalent of the NaSEt may initially be added to the fifth solution to form an intermediate product GaSEtCl₂. A second molar equivalent of the NaSEt may be added to the fifth solution to form another intermediate product Ga(SEt)₂Cl. Finally, a third molar equivalent of the NaSEt may then be added to the fifth solution to form another intermediate product Ga(SEt)₃.

It will be apparent from the above description that, in additional embodiments, instead of using the same reagent (e.g., NaSEt) in each of the sequential additions to the reaction mixture, different reagents including different moieties (other than ethyl groups) may be added to the fourth solution to substitute different, selected moieties into the resulting SSP product at different selected locations within the structure of the SSP molecule or complex.

Example 3

The SSP $(Ph_3P)_2Cu(\mu\text{-SBn})(\mu\text{-SPh})Al(SEt)(SHex)$ may be formed as follows.

The intermediate product $(Ph_3P)_2Cu$ —Cl is formed as described in Example 1 above (and repeated below for convenience). Two (2) molar equivalents of Ph_3P (42.000 grams, 160.13 millimoles) may be added to a mixture of two hundred forty (240) milliliters (mL) of anhydrous benzene (C_6H_6) and two hundred forty (240) milliliters (mL) of anhydrous tetrahydrofuran ((CH $_2$) $_4$ O, THF) to form a first solution, which may be stirred. One (1) molar equivalent of anhydrous Cu(1) Cl (7.9260 grams, 80.064 millimoles) may be added to the stirring first solution to form a first white suspension includ-

ing the intermediate product $(Ph_3P)_2Cu$ —Cl. The first suspension may be concentrated to form a solid product or kept to continue the reaction, which is referred to herein as "Mixture A."

A second solution (which is formed as described above and repeated below for convenience) may be formed by adding one (1) molar equivalent of Na metal (34.000 grams, 1478.9 millimoles, collected from mineral oil and washed several times with diethyl ether) to about 500 mL of diethyl ether. One (1) molar equivalent of EtSH (109.52 mL, 1478.9 millimoles) may be added to the second solution and stirred for twelve (12) hours at forty (40) degrees Celsius (° C.). The EtSH and the Na metal react with one another to form NaSEt. The resulting product that includes the NaSEt (which may comprise a solution or a solid product comprising the NaSEt 15 formed by drying a solution) is referred to herein as "Mixture R."

A third solution (which is formed as described above and repeated below for convenience) may be formed by adding one (1) molar equivalent of Na metal (10.000 grams, 434.97 20 millimoles, collected from mineral oil and washed several times with diethyl ether) to about 200 mL of diethyl ether. One (1) molar equivalent of $C_6H_5CH_2SH(HSBn, 51.066$ mL, 434.97 millimoles) may be added to the third solution and stirred for twelve (12) hours at forty (40) degrees Celsius (° 25 C.). The $C_6H_5CH_2SH$ and the Na metal react with one another to form NaSCH $_2C_6H_5$ (NaSBn). The resulting product that includes the NaSBn (which may comprise a solution or a solid product comprising the NaSBn formed by drying a solution) is referred to herein as "Mixture C."

A fourth solution may be formed by adding one (1) molar equivalent of Na metal (10.000 grams, 434.97 millimoles, collected from mineral oil and washed several times with diethyl ether) to about 200 mL of diethyl ether. One (1) molar equivalent of C_6H_5SH (HSPh, 44.454 mL, 434.97 milli- 35 moles) may be added to the fourth solution and stirred for twelve (12) hours at forty (40) degrees Celsius (° C.). The C_6H_5SH and the Na metal react with one another to form NaSC₆H₅ (NaSPh). The resulting product that includes the NaSPh (which may comprise a solution or a solid product 40 comprising the NaSPh formed by drying a solution) is referred to herein as "Mixture D."

A fifth solution may be formed by adding one (1) molar equivalent of Na metal (10.000 grams, 434.97 millimoles, collected from mineral oil and washed several times with 45 diethyl ether) to about 200 mL of diethyl ether. One (1) molar equivalent of $C_6H_{13}\mathrm{SH}$ (HSHex, 61.227 mL, 434.972 millimoles) may be added to the fifth solution and stirred for twelve (12) hours at forty (40) degrees Celsius (° C.). The $C_6H_{13}\mathrm{SH}$ and the Na metal react with one another to form 50 NaSC $_6H_{13}$ (NaSHex). The resulting product that includes the NaSHex (which may comprise a solution or a solid product comprising the NaSHex formed by drying a solution) is referred to herein as "Mixture E."

A sixth solution may be formed by adding one (1) molar 55 equivalent of NaSBn (8.184 grams, 80.064 millimoles) provided by Mixture C to the first solution ((Ph₃P)₂Cu—Cl) provided by Mixture A, after which the sixth solution may be stirred for twelve (12) hours. The (Ph₃P)₂Cu—Cl and the NaSBn react with one another to form (Ph₃P)₂CuSBn. The 60 resulting mixture may comprise a sixth solution. This solution may comprise the intermediate product (Ph₃P)₂CuSBn (which may comprise a solution or a solid product comprising the (Ph₃P)₂CuSBn formed by drying a solution) and is referred to herein as "Mixture F."

A seventh solution may be formed by adding one (1) molar equivalent anhydrous AlCl₃ (10.676 grams, 80.064 milli-

22

moles) to the sixth solution of $(Ph_3P)_2CuSBn$ provided by Mixture F, after which the seventh solution may be stirred for about sixty (60) minutes at eighty (80) degrees Celsius (° C.). The resulting mixture is the seventh solution. This solution may comprise the intermediate product $(Ph_3P)_2Cu(\mu\text{-SBn})$ ($\mu\text{-Cl}Al(Cl)_2$ (which may comprise a solution or a solid product comprising the $(Ph_3P)_2Cu(\mu\text{-SBn})(\mu\text{-Cl})Al(Cl)_2$ formed by drying a solution) and is referred to herein as "Mixture G."

An eighth solution may be formed by adding one (1) molar equivalent NaSPh (10.581 grams, 80.064 millimoles) provided by Mixture D to the seventh solution of $(Ph_3P)_2Cu(\mu-SBn)(\mu-Cl)Al(Cl)_2$ provided by Mixture G, after which the eighth solution may be stirred for about sixty (60) minutes at eighty (80) degrees Celsius (° C.). The resulting mixture is the eighth solution. This solution may comprise the intermediate product $(Ph_3P)_2Cu(\mu-SBn)$ $(\mu-SPh)Al(Cl)_2$ (which may comprise a solution or a solid product comprising the $(Ph_3P)_2Cu$ $(\mu-SBn)(\mu-SPh)Al(Cl)_2$ formed by drying a solution) and is referred to herein as "Mixture H."

A ninth solution may be formed by adding one (1) molar equivalent NaSEt (6.735 grams, 80.064 millimoles) provided by Mixture B to the eighth solution of $(Ph_3P)_2Cu(\mu\text{-SBn})(\mu\text{-SPh})Al(Cl)_2$ provided by Mixture H, after which the ninth solution may be stirred for about sixty (60) minutes at eighty (80) degrees Celsius (° C.). The resulting mixture is the ninth solution. This solution may comprise the intermediate product $(Ph_3P)_2Cu(\mu\text{-SBn})(\mu\text{-SPh})Al(SEt)(Cl)$ (which may comprise a solution or a solid product comprising the $(Ph_3P)_2Cu(\mu\text{-SBn})(\mu\text{-Ph})Al(SEt)(Cl)$ formed by drying a solution) is referred to herein as "Mixture 1."

A tenth solution may be formed by adding one (1) molar equivalent NaSHex (11.227 grams, 80.064 millimoles) provided by Mixture E to the ninth solution of (Ph₃P)₂Cu(μ-SBn) (μ-SPh)Al(SEt)(Cl) provided by Mixture 1, after which the tenth solution may be stirred for about twelve (12) hours at eighty (80) degrees Celsius (° C.). After stirring, the resulting mixture may comprise a suspension that includes a solution with particles of the SSP (Ph₃P)₂Cu(μ-SBn) (μ-SPh)Al(SEt) (SHex) suspended therein. The SSP may be isolated by filtering of by-product, and the filtrate may be concentrated as necessary or desirable. Optionally, the isolated SSP may be re-crystallized in preparation for using the SSP to form the ternary chalcopyrite material CuAlS₂.

Example 4

The SSP $(Ph_3P)_2Cu(\mu-SePh)_2ln(SePh)_2$ may be formed as follows.

One (1) molar equivalent of the SSP (Ph₃P)₂Cu(μ -SEt)₂ln (SEt)₂ (10.000 grams, 10.555 millimoles), which may be formed as described in Example 1, may be added to about 50 mL of benzene to form a first solution, which may be stirred. Four (4) molar equivalents of C₆H₅SeH (PhSeH, 4.7080 mL, 42.220 millimoles) may be added to the stirring first solution to form a first solution or suspension including the product (Ph₃P)₂Cu(μ -SePh)₂ln(SePh)₂. The first solution or suspension may be concentrated to form a liquid or solid product (or kept to continue the reaction as one pot reaction).

In some embodiments, however, less than four (4) molar equivalents of the PhSeH may initially be added to the first solution. Fractions of the four (4) molar equivalents may be added sequentially to attain various products. For example, one (1) molar equivalent of the PhSeH may initially be added to the first solution to form an intermediate product, the exact nature of which is not known, but is currently believed to be the (Ph₃P)₂Cu(μ-SEt)(μ-SePh)ln(SEt)₂ species. A second molar equivalent of the PhSeH may be added to the first

solution to form another intermediate product, the exact nature of which is not known, but is currently believed to be the $(Ph_3P)_2Cu(\mu\text{-SePh})_2In(SEt)_2$ species. Similarly, a third molar equivalent of the PhSeH may then be added to the first solution to form another intermediate product, the exact 5 nature of which is not known, but is currently believed to be the $(Ph_3P)_2Cu(\mu\text{-SePh})_2In(PhSe)(SEt)$ species. Finally, a fourth molar equivalent of the PhSeH may be added to the first solution to form the final SSP product $(Ph_3P)_2Cu(\mu\text{-SePh})_2In(SePh)_2$.

After all four (4) molar equivalents of the PhSeH have been added to the first solution, the solution may be stirred for an additional one (1) hour. After stirring, the resulting mixture may comprise a solution or suspension that includes a solution with particles of the SSP (Ph₃P)₂Cu(μ -SePh)₂In(SePh)₂ 15 suspended therein. The SSP may be isolated by evaporation of by-product. Optionally, the isolated SSP may be re-crystallized in preparation for using the SSP to form the ternary chalcopyrite material CuInSe₂.

It will be apparent from the above description that, in 20 additional embodiments, instead of using the same reagent (e.g., PhSeH) in each of the sequential additions to the reaction mixture, different reagents including different moieties (other than phenylselenol) may be added to the first solution to substitute different, selected moieties into the resulting 25 SSP product at different selected locations within the structure of the SSP molecule or complex.

In accordance with yet further embodiments of the present invention, polymeric SSPs may be formed by polymerizing SSP molecules or complexes, such as the SSP molecules or 30 complexes formed as previously described herein. For example, an organometallic polymeric SSP of the general formula $[L_2N(ER^1)_a(\mu-ER^2E)_bM'(ER^1)_c(ER^2E)_d]_m$ may be fabricated using Reactions 1 through 3 (which are the same as above and repeated below for convenience) and Reaction 12 35 below:

$$\begin{array}{lll} N-X+2L\rightarrow L_2N-X & Reaction \ 1 \\ \\ L_2N-X+M'X_3\rightarrow {}^{1}\!\!/_2\{L_2N(\mu\!-\!X)_2M'X_2\}_2 & Reaction \ 2 \\ \\ {}^{1}\!\!/_2\{L_2N(\mu\!-\!X)_2M'X_2\}_2+4MER\rightarrow L_2N(\mu\!-\!ER)_2M'(ER)_2 \\ & (+4MX) & Reaction \ 3 \\ \end{array}$$

$$\begin{array}{ll} {\rm L_2N}(\mu\text{-ER})_2M'({\rm ER})_2+n{\rm HE}^1{\rm R}^1{\rm E}^1{\rm H} \to [{\rm L_2N}({\rm ER})_a(\mu\text{-}\\ {\rm E}^1{\rm R}^1{\rm E}^1)_bM'({\rm ER})_c({\rm E}^1{\rm R}^1{\rm E}^1)_d]_m & {\rm Reaction}\ 12 \end{array}$$

wherein L is a Lewis base (that is coordinated to N by a dative bond), each N is individually selected from Group IB atoms, each M is individually selected from Group IA atoms, each M' is individually selected from Group IIIA atoms, each E is individually selected from Group VIA atoms, each E¹ is indi- 50 vidually selected from Group VIA atoms (E and E¹ may be the same or different), each X is individually selected from group VIIA atoms or a nitrate group, n is any number from less than one (1) to four (4) or excess, a is any number from zero (0) to two (2), b is the difference between two (2) and a (b=2-a), c 55 is any number from zero (0) to two (2), d is the difference between two (2) and c (d=2-c), the sum of b and d is equal to n (n=b+d) (if n is four (4) or less), m is any number representing the size of the resulting polymeric molecules or complexes, each of the R groups is individually selected from the 60 group consisting of alkyl, aryl, vinyl, (per)fluoro alkyl, (per) fluoro aryl, silane, and carbamato groups, and each of the R¹ groups is individually selected from the group consisting of α,ω-E¹ functionalized alkyl, vinyl, (per)fluoro alkyl, (per) fluoro aryl, silane, and carbamato groups.

In Reaction 12, up to four (4) units of the ER groups of the $L_2N(\mu\text{-ER})_2M'(ER)_2$ substance may be replaced with $E^1R^1E^1$

24

to give linear or cross-linked polymeric SSPs. In other words, the degree of polymerization may be relatively higher when, for example, n is about four (4), and the degree of polymerization may be relatively lower when, for example, n is about one (1) or less, with intermediate degrees of polymerization being attained when n has an intermediate value between one (1) and four (4). It is noted that the R^1 groups in the four molar equivalents of the $HE^1R^1E^1H$ reagent may differ from one another, and may be individually selected from the group consisting of $\alpha,\omega\text{-}E^1$ functionalized alkyl, aryl, vinyl, (per) fluoro alkyl, (per)fluoro aryl, silane, and carbamato groups. It is noted that the E^1 groups in the four molar equivalents of the $HE^1R^1E^1H$ reagent may differ from one another, and may be individually selected from the Group VIA atoms.

In some embodiments, m may be one (1) or greater than one (1). For example, m may be between one (1) and about 20,000 or higher.

Reactions 1 through 3 may be carried out as previously described herein. Referring to Reaction 12, the non-polymeric organometallic substance $L_2N(\mu-ER)_2M'(ER)_2$ may be polymerized by reacting between about one (1) and about four (4) molar equivalents of $HE^1R^1E^1H$ with the $L_2N(\mu-ER)_2$ M'($ER)_2$ substance to form a polymeric organometallic SSP of the empirical formula $[L_2N(ER)_a(\mu-E^1R^1E^1)_bM'(ER)_c$ ($E^1R^1E^1$)_cd]_m. Reaction 12 may be carried out in a solution comprising benzene (C_6H_6) or a mixture of more than one solvent or another suitable solvent.

Example 5

As a non-limiting example, one (1) molar equivalent of $(Ph_3P)_2Cu(\mu\text{-SEt})_2In(SEt)_2$ (20.000 grams, 21.110 millimoles), which may be formed as previously described in Example 1, may be added to about 100 mL of benzene to form a first solution, which may be stirred. Four (4) molar equivalents of $HSCH_2CH_2SH$ (HSEtSH, 7.0821 mL, 84.438 millimoles) may be added to the stirring first solution to form a first suspension including the product $[(Ph_3P)_2Cu(\mu\text{-SCH}_2CH_2S)_2 In(SCH_2CH_2S)_2]_m$, where m may be between one (1) and about 20,000 or higher. The first suspension may be concentrated to form a solid product (or kept to continue the reaction as in a one pot reaction).

In some embodiments, two or more different SSPs (e.g., containing different N and/or M' atoms) may be polymerized by reacting between about one (1) and about four (4) or excess molar equivalents of HE¹R¹E¹H with the two or more different SSPs to form copolymeric organometallic SSPs of the formula $\{[L_2N^1(ER)_a(\mu-E^1R^1E^1)_bM^{'1}(ER)_a\}$ empirical $(E^{1}R^{1}E^{1})_{d}]_{m}[L_{2}N^{2}(ER)_{e}(\mu - E^{1}R^{1}E^{1})_{e}M^{2}(ER)_{e}(E^{1}R^{1}E^{1})_{e}M^{2}(ER)_{$ E^{1}_{h} _{ln}, wherein L is a Lewis base coordinated to N¹ and N² by a dative bond, each N¹ and N² is individually selected from Group IB atoms (N^1 and N^2 may be the same or different), each M¹ and M² is individually selected from Group IIIA atoms (M¹ and M² may be the same or different), each E and E¹ is individually selected from Group VIA atoms (E and E¹ may be same or different), a is any number from zero (0) to two (2), b is the difference between two (2) and a (b=2-a), c is any number from zero (0) to two (2), d is the difference between two (2) and c (d=2-c), m is any number, e is any number from zero (0) to two (2), f is the difference between two (2) and e (f=2-e), g is any number from zero (0) to two (2), h is the difference between two (2) and g (h=2-g), n is any number, 1 is any number, each R group is individually selected from the group consisting of alkyl, aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl, silane, and carbamato groups, and each of the R¹ groups is individually selected from the group consisting of α,ω -E¹ functionalized alkyl, aryl, vinyl, (per)

fluoro alkyl, (per)fluoro aryl, silane, and carbamato groups. It is noted that the R^1 groups in the four molar equivalents of the $HE^1R^1E^1H$ reagent may differ from one another, and may be individually selected from the group consisting of alkyl, aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl, silane, and carbamato groups. It is noted that the E^1 groups in the four molar equivalents of the $HE^1R^1E^1H$ reagent may be the same or different from one another, and may be individually selected from the Group V1A atoms. It is noted that all possible variations in identities and quantities of different SSPs can be used 10 to form copolymeric single source precursors.

In some embodiments, m, n, and l may be one (1) or greater than one (1). For example, m, n, and l may be between one (1) and about 20,000 or higher (m, n, and l may be the same or different from one another).

Example 6

As a non-limiting example, one (1) molar equivalent of $(Ph_3P)_2Cu(\mu\text{-SEt})_2\ln(\text{SEt})_2$ (10.000 grams, 10.555 milli- 20 moles) and one (1) molar equivalent of $(Ph_3P)_2Cu(\mu\text{-SEt})_2Ga$ (SEt)₂ (9.5240 grams, 10.555 millimoles), which may be provided using methods previously described herein, may be added to about 100 mL of benzene to form a first solution, which may be stirred. Eight (8) molar equivalents of 25 HSCH₂CH₂SH(HSEtSH, 25 10, 25 10 mL, 25 11 mL, 25 11 mL, 25 12 mJ, 25 22 mJ, 25 22 mJ, 25 23 mJ, 25 23 mJ, 25 24 where m, n, and 1 may be between one (1) and about 20,000 or higher. The first suspension may be concentrated to form a solid product (or kept to continue the reaction as in a one pot reaction).

Example 7

As a non-limiting example, one (1) molar equivalent of $(Ph_3P)_2Cu(\mu-SEt)_2ln(SEt)_2$ (10.000 grams, 10.555 millimoles), and one (1) molar equivalent of $(Ph_3P)_2Ag(\mu-SEt)_2$ Ga(SEt)₂ (9.9919 grams, 10.555 millimoles), which may be 40 provided using methods previously described herein, may be added to about 100 mL of benzene to form a first solution, which may be stirred. Eight (8) molar equivalents of HSCH₂CH₂SH (HSEtSH, 7.0821 mL, 84.438 millimoles) may be added to the stirring first solution to form a copolymeric, organometallic SSP having the empirical formula $\{[(Ph_3P)_2Cu(\mu-SCH_2CH_2S)_2ln(SCH_2CH_2S)_2]_m[(Ph_3P)_2Ag(\mu-SCH_2CH_2S)_2Ga(SCH_2CH_2S)_2]_n\}_1$; where m, n, and 1 may be between one (1) and about 20,000 or higher. The first suspension may be concentrated to form a solid product (or 50 kept to continue the reaction as in a one pot reaction).

 $SCH_2CH_2S)_2$ $Ga(SCH_2CH_2S)_2]_m[(Ph_3P)_2Cu(\mu-60)$

SCH₂CH₂S)₂Al(SCH₂CH₂S)₂]_m[(Ph₃P)₂Ag(μ-

SCH₂CH₂S)₂ln(SCH₂CH₂S)₂]_m[(Ph₃P)₂Ag(µ-SCH₂CH₂S)₂ Ga(SCH₂CH₂S)₂]_m[(Ph₃P)₂Ag(µ-SCH₂CH₂S)₃]_m[(Ph₃P)₂Ag(µ-SCH₂CH₂S)₂]_m[(Ph₃P)₂Ag(µ-SCH₂CH₂S)₂]_m[(Ph₃P)₂Ag(µ-SCH₂CH₂S)₂]_m[(Ph₃P)₂Ag(µ-SCH₂CH₂S)₂]_m[(Ph₃P)₂Ag(µ-SCH₂CH₂S)₂]_m[(Ph₃P)₂Ag(µ-SCH₂CH

 $SCH_2CH_2CH_2S)_2$ $Im\{(H_3F)_2Ag(\mu^2SCH_2CH_2S)_2\}_m\}_m$, wherein each m may be between one (1) and about 20,000 or higher.

In some embodiments, two or more different $HE^1R^1E^1H$ species may be reacted with one or more than one $L_2N(\mu$ -

26

ER)₂ M'(ER)₂ species in a polymerization reaction to form a copolymeric organometallic SSP having the empirical formula $\{[L_2N^1(\mu-ER)_a(\mu-E^1R^1E^1)_bM^1(ER)_c(E^1R^1E^1)_d]_m \{L_2N^2(\mu-ER)_e(\mu-E^1R^1E^1)_dM^2(ER)_e(E^1R^1E^1)_h]_n\}_{I}$

In some embodiments, one or more different $HE^1R^1E^1H$ species and one or more different HE^2R^2 species may be reacted with one or more different $L_2N(\mu\text{-ER})_2M'(ER)_2$ species to form a polymeric organometallic SSP of the empirical formula $\{[L_2N(E^2R^2)_a(\mu\text{-}E^1R^1E^1)_bM'(E^2R^2)_c(E^1R^1E^1)_d\}_m$. It is noted that the E, E^1 , and E^2 atoms in the reagents may be the same or different, and may be individually selected from the Group V1A atoms.

Example 8

As a non-limiting example, one (1) molar equivalent of (Ph₃P)₂Cu(μ-SEt)₂ln(SEt)₂ (10.000 grams, 10.555 millimoles), which may be provided using methods previously described herein, may be added to about 100 mL of benzene to form a first solution, which may be stirred. Four (4) molar equivalent of C₆H₅SeH (PhSeH, 4.4837 mL, 42.220 millimoles) and one (1) molar equivalent of HSCH₂CH₂SH (HSEtSH, 0.88530 mL, 10.555 millimoles) may be added to the stirring first solution to form copolymeric organometallic SSPs having the empirical formula [(Ph₃P)₂Cu(μ-SCH₂CH₂S)(μ-SePh)ln(SePh)₂]_m or [(Ph₃P)₂Cu(μ-SePh)₂ln (SePh)(SCH₂CH₂S)]_m, where m may be between one (1) and about 20,000 or higher. The first suspension may be concentrated to form a solid product (or kept to continue the reaction as in a one pot reaction).

In accordance with additional embodiments of the present invention, a polymeric SSP of the general formula $[L_2N(ER)_a (\mu-E^1R^1E^1)_bM'(ER)_c(E^1R^1E^1)d]_m$ may be fabricated using Reactions 1 and 2 (which are the same as above and repeated below for convenience), and additional Reaction 13 below:

$$N - X + 2L \rightarrow L_2N - X$$
 Reaction 1

$$L_2N$$
— $X+M'X_3 \rightarrow \frac{1}{2} \{L_2N(\mu-X)_2M'X_2\}_2$ Reaction 2

$$\begin{array}{ll} ^{1/2} \{L_{2}N(\mu - X)_{2}M'X_{2}\}_{2} \rightarrow nME^{1}R^{1}E^{1}M + pMER \rightarrow [L_{2}N\\ (ER)_{a}(\mu - E^{1}R^{1}E^{1})_{b}M'(ER)_{c}(E^{1}R^{1}E^{1})_{d}]_{m}(+4MX) & \text{Reaction } 13 \end{array}$$

wherein L is a Lewis base that is coordinated to N by a dative bond, each N is individually selected from Group 1B atoms, each M is individually selected from Group 1A atoms, each M' is individually selected from Group 111A atoms, each X is individually selected from Group VIIA atoms or a nitrate group, each E and E¹ is individually selected from Group V1A atoms (E and E^1 may be the same or different), n is any number from one (1) to four (4), p is the difference between four (4) and n (p=4-n), a is any number from zero (0) to two (2), b is the difference between two (2) and a (b=2-a), c is any number from zero (0) to two (2), d is the difference between two (2) and c (d=2-c), the sum of b and d is equal to n (n=b+d), the sum of a and c is equal top (p=a+c), m is any number representing the size of the resulting polymeric molecules or complexes, each of the R groups is individually selected from the group consisting of alkyl, aryl, vinyl, (per) fluoro alkyl, (per)fluoro aryl, silane, and carbamato groups, and each of the R1 groups is individually selected from the group consisting of α, ω -E¹ functionalized alkyl, aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl, silane, and carbamato groups. In some embodiments, m may be one (1) or greater than one (1). For example, m may be between one (1) and about 20,000 or greater.

Reactions 1 and 2 may be carried out as previously described herein. Referring to Reaction 13, the non-poly-

meric organometallic substance $L_2N(\mu-X)_2M'(X)_2$ may be polymerized by reacting between about one (1) and about four (4) molar equivalents of $ME^1R^1E^1M$ and MER (either sequentially or simultaneously) with the $L_2N(\mu-X)_2M'(X)_2$ substance to form a polymeric organometallic SSP of the empirical formula $[L_2N(ER)_a(\mu-E^1R^1E^1)_bM'(ER)_c(E^1R^1E^1)_d]_m$. Reaction 13 may be carried out in a solution comprising benzene (C_6H_6) or another suitable solvent or solvent mixture.

Example 9

As a non-limiting example, one (1) molar equivalent of $\frac{1}{2}\{(Ph_3P)_2Cu(\mu-Cl)_2ln(Cl)_2\}_2$ (67.635 grams, 80.064 millimoles), which may be provided using methods previously described herein, may be added to a mixture of about 240 mL of anhydrous benzene and 240 mL of anhydrous THF to form a first solution, which may be stirred. About two (2) molar equivalents of NaSCH_2CH_2SNa (22.123 grams, 160.128 millimoles) and about two (2) molar equivalents of NaSEt (13.470 mL, 160.128 millimoles) may be added to the stirring first solution to form a polymeric, organometallic SSP having the empirical formula $[(Ph_3P)_2Cu(\mu-(SCH_2CH_2S))(\mu-SEt)]_m$, where m may be between one (1) and about 20,000 or higher.

In some embodiments, two or more different $L_2N(\mu-X)_2M'$ 25 (X)₂ species may be polymerized by reacting between about one (1) and about four (4) molar equivalents of ME¹R¹E¹M and MER (either sequentially or simultaneously) with the two or more different $L_2N(\mu-X)_2M'(X)_2$ species to form a copolymeric organometallic SSP of the empirical formula $\{[\hat{L}_2N^1]^30\}$ $(ER)_a(\mu-E^1R^1E^1)_bM^{'1}(ER)_c(E^1R^1E^1)_d]_m[L_2N^2(ER)_e(\mu-E^1R^1E^1)_d]_m$ $E^1R^1E^1$)_A M^2 (ER)_g($E^1R^1E^1$)_h]_n}_t, wherein \tilde{L} is a Lewis base that is coordinated to N^1 and N^2 by dative bonds, each N^1 and N² is individually selected from Group 1B atoms (N¹ and N² may be the same or different), each M is individually selected 35 from Group 1A atoms, each M¹ and M² is individually selected from Group 111A atoms (M¹ and M² may be the same or different), each E and E¹ is individually selected from Group V1A atoms (E and E^1 may be the same or different), each X is individually selected from Group V11A atoms or a 40 nitrate group, a is any number from zero (0) to two (2), b is the difference between two (2) and a (b=2-a), c is any number from zero (0) to two (2), d is the difference between two (2) and c (d=2-c), m is any number, e is any number from zero (0) to two (2), f is the difference between two (2) and e (f=2-e), 45 g is any number from zero (0) to two (2), h is the difference between two (2) and g (h=2-g), n is any number, 1 is any number, each of the R groups is individually selected from the group consisting of alkyl, aryl, vinyl, (per)fluoro alkyl, (per) fluoro aryl, silane, and carbamato groups, and each of the R¹ 50 groups is individually selected from the group consisting of α,ω -E¹ functionalized alkyl, aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl, silane, and carbamato groups. It is noted that the E^1 groups in the four molar equivalents of the $ME^1R^1E^1M$ reagent may differ from one another, and may be individually 55 selected from the Group VIA atoms. It is noted that all possible variations in identities and quantities of any organometallic substance can be used to form copolymeric single source precursors. In some embodiments, m, n, and I may be one (1) or greater than one (1). For example, m, n, and 1 may 60 be between one (1) and about 20,000 or higher (m, n, and 1 may be the same or different).

Example 10

As a non-limiting example, one (1) molar equivalent of $\frac{1}{2}\{(Ph_3P)_2Cu(\mu-Cl)_2ln(Cl)_2\}_2$ (33.818 grams, 40.032 milli-

28

moles) and one (1) molar equivalent of ½{(Ph₃P)₂Cu(μ-Cl)₂ $Ga(Cl)_2$ ₂ (32.012 grams, 40.032 millimoles), which may be formed using methods previously describe herein, may be added to a mixture of about 240 mL of anhydrous benzene and 240 mL of anhydrous THF to form a first solution, which may be stirred. About two (2) molar equivalents of NaSCH₂CH₂SNa (22.123 grams, 160.128 millimoles) and about two (2) molar equivalents of NaSEt (13.470 mL, 160.128 millimoles) may be added to the stirring first solution 10 to form a copolymeric, organometallic SSP having the empirical equation $\{[(Ph_3P)_2Cu(\mu-(SCH_2CH_2S))(\mu-SEt)]$ ln $(SCH_2CH_2S)(SEt)]_m[(Ph_3P)_2Cu(\mu-(SCH_2CH_2S))(\mu-SEt)G$ $a(SCH_2CH_2S)(SEt)]_n$ _l, where m, n, and 1 may be between one (1) and about 20,000 or higher. The first suspension may be concentrated to form a solid product (or kept to continue the reaction as one pot reaction).

Example 11

As a non-limiting example, one (1) molar equivalent of $\frac{1}{2}$ {(Ph₃P)₂Cu(μ -Cl)₂ln(Cl)₂}₂ (33.818 grams, 40.032 millimoles) and one (1) molar equivalent of $\frac{1}{2}$ {(Ph₃P)₂Cu(μ -Cl) $_2$ Ga(Cl) $_2$ $\}_2$ (32.012 grams, 40.032 millimoles), which may be provided using methods previous described herein, may be added to a mixture of about 240 mL of anhydrous benzene and 240 mL of anhydrous THF to form a first solution, which may be stirred. About four (4) molar equivalents of NaSCH₂CH₂SNa (NaSEtSNa, 44.246 grams, 320.256 millimoles) may be added to the stirring first solution to form a copolymeric, organometallic SSP having the empirical formula $\{[(Ph_3P)_2Cu(\mu-(SEtS))_2]n(SEtS)_2\}_m[(Ph_3P)_2Cu(\mu-(SEtS))_2]$ $(SEtS)_2Ga(SEtS)_2]_n$, where m, n, and 1 may be between one (1) and about 20,000 or higher. The first suspension may be concentrated to form a solid product (or kept to continue the reaction as a one pot reaction).

In some embodiments, two or more different $ME^1R^1E^1M$ and MER species may be polymerized by reacting between one or more than one of the $L_2N(\mu-X)_2M'(X)_2$ species to form a copolymeric organometallic SSP of the empirical formula $\{[L_2N^1(ER)_a(\mu-E^1R^1E^1)_bM'^1(ER)_c(E^1R^1E^1)_d]_m[L_2N^2(ER)_e(\mu-E^1R^1E^1)_dM'^2(ER)_g(E^1R^1E^1)_h]_n\}_I$.

Using the polymerization reactions described above, new polymeric, organometallic SSPs having the empirical formula $[L_2N(ER)_a(\mu-E^1R^1E^1)_bM'(ER)_c(E^1R^1E^1)_d]_m$ may be provided, wherein each of a, b, c, and d are between zero (0) and two (2). For example, in some embodiments, the SSPs may have the empirical formula $[L_2N(\mu-ER)(\mu-E^1R^1E^1)M'(ER)_c(E^1R^1E^1)_d]_m$. In additional embodiments, the SSPs may have the empirical formula $[L_2N(\mu-E^1R^1E^1)_2M'(ER)_c(E^1R^1E^1)_d]_m$. In additional embodiments, the SSPs may have the empirical formula $[L_2N(ER)_a(\mu-E^1R^1E^1)_bM'(ER)_c(E^1R^1E^1)]_m$. In yet further embodiments, the SSPs may have the empirical formula $[L_2N(ER)_a(\mu-E^1R^1E^1)_bM'(E^1R^1E^1)_d]_m$.

In some embodiments, the polymeric, organometallic SSPs may comprise substantially linear polymeric molecules or complexes. In additional embodiments, the polymeric, organometallic SSPs may comprise cross-linked substantially linear polymeric molecules or complexes. In yet further embodiments, the cross-linking between the individual SSP molecules or complexes may be so extensive that the polymeric, organometallic SSPs have a three-dimensional network structure.

The reaction pathways disclosed herein for forming SSPs allow the different R groups in the SSPs, as well as the different E, N, and M' atoms, to be selectively and individually tailored, which further allows chemical and physical

properties of the SSPs, such as, for example, reactivity, solubility, melting point, boiling point, etc., to be selectively

SSPs (including polymeric SSPs and non-polymeric SSPs) made in accordance with embodiments of the present invention may be used to form chalocogenide materials (e.g., semiconductive ternary chalcopyrite materials) using methods such as, for example, those disclosed in U.S. patent application Ser. No. 12/047,956, filed Mar. 13, 2008, now U.S. Pat. No. 8,003,070, issued Aug. 23, 2011 to Fox et al., the disclosure of which is incorporated herein in its entirety by this reference.

Semiconductor materials and particles fabricated using SSPs formulated in accordance with embodiments of methods of the present invention may be used in many different 15 types of semiconductor devices. For example, semiconductor materials and particles formed using embodiments of methods of the present invention may be used in semiconductor devices such as, for example, diodes (e.g., light emitting diodes (LEDs)), photovoltaic devices, sensors, solid state 20 lasers, and integrated circuits (e.g., memory modules and microprocessors).

While the invention may be susceptible to various modifications and alternative forms, specific embodiments have detail herein. However, it should be understood that the invention is not intended to be limited to the particular forms disclosed. Rather, the invention includes all modifications, equivalents, and alternatives falling within the scope of the their legal equivalents.

What is claimed is:

- 1. A method of forming a single-source precursor for use in forming a chalcogenide material, comprising:
 - mula $\frac{1}{2}\{L_2N(\mu-X)_2M'X_2\}_2$; and
 - reacting MER with the intermediate product to form a single source precursor having the empirical formula $L_2N(\mu-ER)_2M'(ER)_2;$
 - wherein L is a Lewis base coordinated to N by a dative 40 bond, each N is individually selected from Group IB atoms, each M is individually selected from Group IA atoms, each M' is individually selected from Group IIIA atoms, each E is individually selected from Group VIA atoms, each X is individually selected from Group VIIA 45 atoms or a nitrate group, and each R group is individually selected from the group consisting of alkyl, aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl, silane, and carbamato groups; and
 - wherein reacting MER with the intermediate product com- 50 prises reacting each of a first MER species and a different second MER species with the intermediate product such that at least one E differs from at least one other E or at least one R group differs from at least one other R group in the single source precursor.
- 2. The method of claim 1, further comprising selecting each of the first MER species and the different second MER

30

species such that at least one R group in the single source precursor differs from at least one other R group in the single

- 3. The method of claim 1, further comprising selecting the Lewis base to comprise a substituted phosphine.
- 4. The method of claim 1, wherein reacting MER with the intermediate product to form the single source precursor comprises reacting less than four molar equivalents of first MER species with one molar equivalent of the intermediate product to form another intermediate product, and subsequently reacting an additional quantity of second MER species with the another intermediate product to form the single source precursor.
- 5. The method of claim 4, further comprising forming the another intermediate product to have an empirical formula selected from the group consisting of $\frac{1}{2}\{L_2N(\mu-X)(\mu-ER)\}$ $(ER)_2$ ₂, and $\frac{1}{2}\{L_2N(\mu-X)(\mu-ER)M'(ER)(X)\}_2$.
- 6. The method of claim 4, further comprising selecting the R group of the additional quantity of second MER species to differ from the R group of the less than four molar equivalents of first MER species.
- 7. The method of claim 6, further comprising selecting the been shown by way of example and have been described in 25 E of the additional quantity of second MER species to differ from the E of the less than four molar equivalents of first MER species.
- 8. The method of claim 4, further comprising selecting the E of the additional quantity of second MER species to differ invention as defined by the following appended claims and 30 from the E of the less than four molar equivalents of first MER species.
 - 9. The method of claim 1, wherein at least one R group comprises an ethyl group.
 - 10. The method of claim 1, further comprising isolating the forming an intermediate product having the empirical for- 35 intermediate product prior to reacting MER with the intermediate product.
 - 11. A method of forming a single-source precursor for use in forming a chalcogenide material, comprising:
 - forming an intermediate product having the empirical formula $\frac{1}{2}\{L_2N(\mu-X)_2M'X_2\}_2$; and
 - reacting MER with the intermediate product to form a single source precursor having the empirical formula $L_2N(\mu-ER)_2M'(ER)_2;$
 - wherein L is a Lewis base coordinated to N by a dative bond, each N is individually selected from Group IB atoms, each M is individually selected from Group IA atoms, each M' is individually selected from Group IIIA atoms, each E is individually selected from Group VIA atoms, each X is individually selected from Group VIIA atoms or a nitrate group, and each R group is individually selected from the group consisting of alkyl, aryl, vinyl, (per)fluoro alkyl, (per)fluoro aryl, silane, and carbamato groups; and
 - wherein the intermediate product is isolated prior to reacting MER with the intermediate product.