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(NASA-Case-XLE-02545-1) METHOD FOR THE
PREPARATION OF INORGANIC SINGLE CRYSTAL AND
POLYCRYSTALLINE ELECTRONIC MATERIALS Patent
(NASA) 3 p

TO: XXX/Scientific & Technical Information Division
Attn: Miss Winnie M. Morgan

FROM: GP/Office of Assistant General
Counsel for Patent Matters

SUBJECT: Announcement of NASA-Owned U.S. Patents in STAR

In accordance with the procedures agreed upon by Code GP and Code
KSI, the attached NASA-owned U.S. Patent is being forwarded for
abstracting and announcement in NASA STAR.

The following information is provided:

U.S. Patent No. : 3,429,756
Government or Corporate Employee : Monsanto Chem. Prod. Co.
Supplementary Corporate Source (if applicable) :
NASA Patent Case No. : W-245 XLE-2545

NOTE - If this patent covers an invention made by a corporate employee of a NASA Contractor, the following is applicable:

YES [ ] NO [ ]

Pursuant to Section 305(a) of the National Aeronautics and Space Act, the name of the Administrator of NASA appears on the first page of the patent; however, the name of the actual inventor (author) appears at the heading of column No. 1 of the Specification, following the words "...with respect to an invention of ..."

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Enclosure
METHOD FOR THE PREPARATION OF INORGANIC SINGLE CRYSTAL AND POLYCRYSTALINE ELECTRONIC MATERIALS

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4 Claims

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The invention described herein was made in the performance of work under a NASA contract and is subject to the provisions of the National Aeronautics and Space Act of 1958, Public Law 85–568 (72 Stat. 426; 42 U.S.C. 2451) as amended.

This invention relates to a method for producing self-supporting semiconductor crystals suitable for device fabrication. More particularly, this invention relates to a method for producing semiconductor crystals of large area and having smooth, flat, damage-free surfaces. Still more particularly this invention involves the removal of substrate materials, by selective etching, from epitaxial overgrowths of semiconductor materials deposited thereon to expose a smooth, flat, damage-free surface of said film suitable for device fabrication. The epitaxial deposits herein are thick enough to be self-supporting.

In the fabrication of electronic devices it is highly desirable and frequently imperative that the semiconductor component thereof be essentially free of surface defects and work damage. It is also necessary that the thin film semiconductor component be smooth, flat and of uniform thickness for optimum performance.

The production and reliable utilization of many electronic devices having the requisite electrical properties for various device applications has been hindered by lack of single crystal semiconductor components having the above physical characteristics. Various methods are known and have been used to deposit single crystal semiconductor materials from a vapor phase. The more practical methods involve vapor deposition of single crystal materials epitaxially upon seed crystals of the same or different material. The substrate frequently serves merely to support the epitaxial overgrowth. Among the methods used for epitaxial deposition of single crystal materials might be mentioned the use of various transport agents to carry the semiconductor component or compound from a source region to a deposition region. Examples of such transport systems are the use of elemental halogens, halogen compounds or hydrogen halides as transport agents carried in inert gases, e.g., hydrogen, helium or argon. Other vapor deposition methods involve thermal decomposition or hydride reduction of compounds containing semiconductor elements which react and deposit in single crystal form on a seed crystal substrate.

In all of the foregoing vapor deposition processes, rigid control must be exercised over process conditions such as substrate preparation and seed crystal substrates, maintaining proper reaction conditions such as flow rates of reactants, relative concentrations of carrier gases to reactants, temperature gradients in the deposition zone, etc. Reactant materials often must be very pure such as in the preparation of GaP from its elements carried by H2 in HCl.

The use of etchants to remove surface defects often creates other problems; e.g., it is very difficult, if not impossible, to control the degree and depth of penetration of the etchant into the material, hence, the semiconductor crystal frequently is of non-uniform thickness; also, the surface frequently has an irregular contour or wavy appearance and the crystal frequently has rounded corners and edges.

Another approach to the problem of obtaining smooth, flat single crystal materials having large area for device fabrication involves the vapor deposition of epitaxial overgrowth on various substrates as described above. Subsequently, the substrate material is removed by lapping to expose the surface of the single crystal epitaxial layer contiguous to and forming an interface with said substrate. When the deposition procedure has been carefully conducted on substrate crystals the surface of which has been properly prepared, the junction between the epitaxial overgrowth and substrate crystal will be well-defined, smooth and flat, having large contact area. Therefore, by removing the substrate the thin layer of epitaxial material contiguous thereto is made available for device fabrication. However, the removal of the substrate layer by mechanical means as by sawing and/or lapping subjects the epitaxial crystal to work damage effects as described above. Moreover, it is most difficult to remove the substrate by lapping in such manner that the lapped surface of the substrate is parallel to surface of the overgrowth in contact therewith. Hence, by this operation the entire interface surface of the epitaxial layer is not exposed and made available for device fabrication. Some of the overgrowth material must then be removed to compensate for this non-parallelism and expose the entire surface of the epitaxial layer.

Accordingly, it is an object of this invention to provide self-supporting semiconductor materials suitable for device fabrication. Another object of this invention is to provide single crystal semiconductor crystals which are smooth, flat, essentially free of surface defects and work damage and have relatively large area.

Still another object of the invention is to provide a method for obtaining the above described semiconductor materials which is simple, rapid and economical.

Yet another object of the invention is the provision of a method for obtaining single crystal semiconductor materials as described above which is not dependent upon any particular method of preparing and depositing such single crystals.

Another object of this invention is to provide a method for obtaining the desired single crystal material which substantially eliminates substrate removal by lapping with its attendant handicaps as described above. According to the present invention it has been discovered that smooth, flat, damage-free surfaces of large area can be made available for device fabrication by the use of a selective etching operation which completely removes the substrate upon which the desired single crystal material has been deposited with negligible
effect on the single crystal material itself which is substantially impermeable to the reagents used.

In a preferred embodiment the use of selective oxidizing etchants in which the solution rates of the substrate materials is high relative to the solution rate of the epitaxial material deposited thereon. Preferably, the solution rate of the epitaxial material will always be low and the higher the ratio of these respective solution rates, i.e., solution rate of substrate relative to solution rate of the epitaxial film, in a given etchant, the better.

The various etchants contemplated herein include both acidic and basic etching solutions which are polishing etches relative to the material. Exemplary etchants include mixtures of alkali metal hydroxides, e.g., lithium hydroxide, sodium hydroxide, potassium hydroxide, and alkalis e.g., methanol, propanol, butanol and the like; concentrated nitric acid; mixtures of hydrofluoric acid, nitric acid and water with or without acetic acid; boric oxide and lead oxide; perchloric acid; sodium carbonate, ammonium chloride; various salts such as sodium fluoride. Various mixtures of these etchants are also contemplated.

The relative proportions of the ingredients in etching mixtures and etching temperatures can be varied over a wide range and thereby control etching times. In general, semiconductor crystals prepared according to the present invention contemplate inorganic single crystal semiconductor materials such as silicon, germanium, compounds and alloys of elements selected from Groups II and VI (II-VI compounds) and elements from Groups III and V (III-V compounds) of Mendeleev's Periodic System. These semiconductor single crystals preferably are epitaxially deposited on substrates of a dissimilar semiconductor material, thus forming heterojunctions. However, substrates of the same material as the epitaxial overgrowth (homojunctions) are also contemplated wherein the substrate or epitaxial overgrowth is so doped as to induce in one material, e.g., heavily doped n-type GaAs, significantly different solution rate with respect to the other material, e.g., lightly doped n-type GaAs. It is also within the purview of this invention to prepare organic semiconductor single crystals e.g., metal polyphthalocyanines, particularly copper polyphthalocyanine, in the same manner for device fabrication. In addition, it is within the purview of this invention to obtain polycrystalline materials suitable for use in various electronic devices.

In broad aspect the present invention is applicable to the preparation of semiconductor materials, particularly single crystal materials, for use in devices which depend on electrical properties of a thin film surface layer. As mentioned above, preferred materials herein include single crystals of silicon, germanium, II-VI compounds and alloys such as the sulfides, selenides and tellurides of zinc, cadmium and mercury and III-V compounds such as the nitrides, phosphides, arsenides and antimonides of bismuth, aluminum, gallium and indium. Among the electronic devices which utilize the single crystal materials produced herein are transistors, rectifiers, varactors, diodes, switching devices, photoelectric cells such as photoconductors, photovoltaic cells, e.g., solar cells, electroluminescent devices, detectors, resistors, amplifiers and the like.

The invention will be better understood by reference to the specific embodiments present below.

EXAMPLE 1

This example illustrates the preparation and recovery of single crystal gallium phosphide, GaP.

Into a fused silica reaction tube was placed a small quantity of red phosphorus near the source end of the tube, elemental gallium near the center of the tube and a seed crystal of gallium arsenide near the deposition end of the tube. The fused silica tube was then placed into three furnaces as indicated earlier, the source GaP was heated to 890° C. Phosphorus was carried over the gallium and gallium arsenide heated to about 1000° C. for about 20 minutes. The gallium and gallium arsenide temperatures were then reduced to about 890° C. and 815° C., respectively. Hydrogen chloride was then admixed with the hydrogen and this mixture carrying vaporized phosphorus passed over the gallium source into the deposition zone, containing the gallium arsenide seed crystal.

Flow rates for the HCl:H2-P2 mixture were 150:1:33 cc./min. Gallium phosphide was deposited as an epitaxial film on the substrate. The interface between the epitaxial film and substrate was smooth and flat and the upper surface of the film contained microscopically rough mat surface having some small nodular growths.

This surface was further treated by passing phosphorus vapor over hydrogen but without HCl over the GaP/GaAs structure heated to 1000° C. Again, the source and substrate temperatures were reduced and HCl introduced to deposit more GaP. This procedure was repeated five times, thus building up the thickness of the overgrowth to about 175 microns. The surface of the GaP is somewhat irregular due to a build up of the nodular growths after repeated depositions.

The GaAs substrate is completely dissolved in the nitric acid which does not dissolve the GaP. The bottom surface of the GaP, now exposed after the nitric acid treatment is essentially smooth, flat and damage free. This nitric acid etching treatment lasted about 16 hours.

EXAMPLE 2

In an alternative embodiment, GaP was used as the source material instead of elemental gallium and phosphorus, although the phosphorus-hydrogen pretreatment of the GaAs substrate was retained. Hydrogen chloride and hydrogen, at flow rates of 1:150 cc./min., respectively, were passed over the source GaP heated to 890° C. The GaP was carried to the deposition zone heated to about 850° C. where GaP deposited as an epitaxial film on the GaAs substrate. Again, the GaP/GaAs structure was immersed in hot concentrated nitric acid for about 15 minutes to selectively etch away the GaAs substrate, leaving a single crystal of GaP having a smooth, flat and damage free surface, on which only slight traces of an etch pattern were seen.

EXAMPLE 3

In this example n+ GaAs is used as substrate and n-type GaAs is epitaxially deposited thereon in the manner described in the preceding example. Tellurium is used as dopant in the n+ GaAs in a concentration 1 X 10^15 atoms/cm.
cc. Upon completion of the reaction the n+ GaAs/n-GaAs structure is immersed in an etching solution of HF:HNO3: H2O in the ratio of 1:3:4 parts by volume, for about 10 hours. The n+ GaAs is selectively dissolved in the etchant leaving a single crystal of n-type GaAs with an essentially smooth, flat and damage free surface.

EXAMPLE 4

In this example single crystal zinc sulfide is deposited as an epitaxial film on a substrate of cadmium telluride in the manner described in Example 2. Upon completion of the reaction the zinc CdTe structure is immersed in an etching solution of potassium hydroxide saturated with chlorine for about one hour. The substrate cadmium sulfide is selectively dissolved in the etchant leaving a single crystal of zinc sulfide, the interface surface of which is essentially smooth, flat, uniform and damage free.

The foregoing embodiments are fully illustrative of the invention described herein and are not intended to be limiting of the various modifications that
will occur to those skilled in the art having the present disclosure to guide them.

1. Process for the preparation of semiconductor single crystals having a substantially smooth, flat, uniform and damage-free surface, and being selected from the group consisting of III-V compounds and alloys thereof which comprises immersing an epitaxial film of one of said single crystals deposited on a compatible substrate material which is selected from the same group as said single crystals, in an etching solution in which said substrate material has a higher solution rate than said epitaxial single crystal and is selectively dissolved and said epitaxial single crystal is essentially unaffected.

3. Process according to claim 2 wherein said etching solution is selected from the group consisting of oxidizing acids and bases and salts.

4. Process according to claim 3 wherein said etching solution comprises concentrated nitric acid.

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J. STEINBERG, Primary Examiner.

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