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Module 01: Junction Analysis

19/7/2014

PN Junction diode: Basic structure, Energy band diagrams, zero applied bias, Forward Applied Bias, Reverse applied bias, PN Junction current, small signal model of PN Junction, Generation and Recombination of currents, junction breakdown

• Story So Far :

1) We calculated e^- & h^+ conc's in thermal equilibrium and determined the position of the Fermi level.

2) We then considered the nonequilibrium condition in which excess e^- s and h^+ are present in the semiconductor.

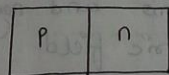
We wish to consider the situation in which a p-type and an n-type semiconductor are brought into contact with one another to form a pn junction.

Analysis of pn junction establishes some basic terminology and concepts that are used in the discussion of other SC devices.

• Understanding the physics of the pn junction is, therefore, an important step in the study of SC devices.

• Basic structure of PN Junction: -
(Homojunction)

Fig: Simplified geometry of a pn junction



↑ Metallurgical junction

↳ (Interface separating n and p regions)

Note: Entire semiconductor is a single-crystal material in which one region is doped with acceptor impurity atoms to form the p-region and n-region is formed by doping it with donor impurity atoms.

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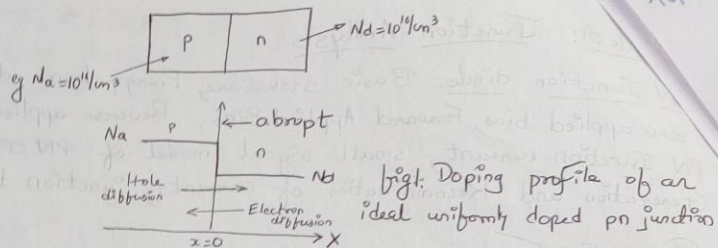


fig1: Doping profile of an ideal uniformly doped pn junction

Analysis of PN-Junction: (Basic structure)

- Consider a step junction wherein we assume
 - Uniform doping in each region.
 - Abupt change in doping at the junction. (Abupt junction approximation) i.e. Doping changes abruptly from p-side to n-side.
- Junction has a very large density gradient in e^- and hole concentrations.
- \therefore Majority carrier e^- s in the n-region starts diffusing into p-region \Rightarrow leaving behind +vely charged donor atoms.
- Also, Majority carrier holes in the p-region diffuse into n-region \Rightarrow leaving behind -vely charged acceptor atoms.
- Positive charge due to donor atoms and negative charge due to acceptor atoms induce an Electric field in the region near the junction.
- The E-field is directed from n to p region.

Assumptions:
1. Boundary between space-charge and neutral region is abrupt.

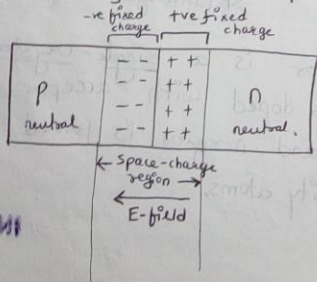


fig2: Space-charge region, electric field for a pn junction.

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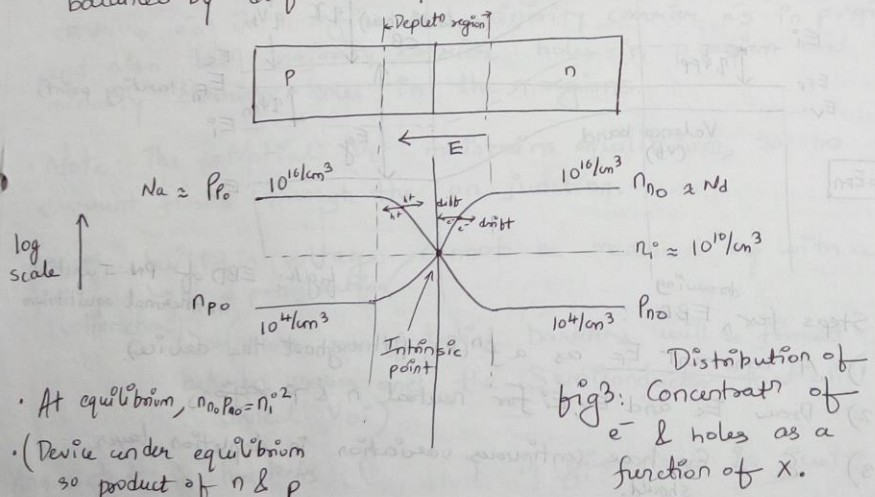
Semiconductor band
 Fermi-Dirac fraction
 physical interpretation of Fermi
 position of n_i with E_f
 Sources of carriers
 Charge-balance

Junction Analysis

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Space-charge region does not contain free e^- and holes, \therefore it is called "depletion region." (coz it is depleted of any mobile carriers)
 coz (essentially all e^- and holes are swept out of the space-charge region by the E-field)

- A density gradient exists in the majority carrier concⁿ at the edges of space-charge region, which produces a diffusion force that pushes the majority carriers towards the space-charge region.
- The built-in E-field in the space-charge region produces "drift force" on the e^- s and holes in the direction opposite to "diffusion force".
- In thermal equilibrium conditions, diffusion is exactly balanced by drift tendency.



Distribution of fig 3: Concentration of e^- & holes as a function of X.

- At equilibrium, $n_0 p_0 = n_i^2$
- (Device under equilibrium so product of n & p should be n_i^2 everywhere).
- Note: Even though doping - changes abruptly at Junction, concentration of free e^- s and holes cannot change abruptly, rather it changes gradually, as shown in fig 3.