## TRAnslational research in Clinical Oncology (TRACO)

### **Program Director**

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### SYLLABUS

| DATE     | TOPIC  |
|----------|--|
|          | SPEAKERS   |
| Sept. 6  | Introduction, Tumor Imaging Moody, Choyke        |
| Sept. 12 | Prostate cancer, Immuno Checkpoint Madan, Goff   |
| Sept. 19 | Lymphoma, Ovarian cancer<br>Dunleavy, Annunziata |
| Sept. 26 | Clinical Trials, HIV                             |
|          | Smith, Maldarelli                                |
| Oct. 3   | TGFbeta, Topoisomerase                           |
|          | Jakowlew, Pommier                                |
| Oct. 10  | COLUMBUS DAY HOLIDAY                             |

### SYLLABUS, continued

| DATE    | TOPIC SPEAKERS  |
|---------|---|
| Oct. 12 | Radiation oncology, Small molecules Nichols, Simeonov |
| Oct. 17 | Epidemiology, SCLC<br>Caporaso, Chen                  |
| Oct. 24 | Breast cancer, Cancer health disparities Zia, Ryan    |
| Oct. 31 | Genomics, Case reports<br>Wei, Olaku                  |
| Nov. 7  | NSCLC, Cervical cancer<br>Szabo, Schiller             |

### SYLLABUS, continued

**TOPIC** DATE **SPEAKERS** Nov. 14 **Functional Genomics, HSP90** Caplen, Neckers Nov. 21 **Epigenetics, Pancreatic cancer** Verma, Hussain Precision medicine, Nanotechnology Nov. 28 Harris, Dobrovolskaia,

#### REGISTRATION

The course is open to all interested personnel without charge.
Registration is available at the NCI CCR Web site

(http://ccr.cancer.gov/trainee-resources-courses-workshops-traco/)

### **CCR** component

Registrants can attend tumor boards, grand rounds, visit technology and/or core facilities. Please contact Dr. Moody, if interested to make appropriate reservations.

### COURSE CERTIFICATION

Registrants can obtain a course certificate upon passing a computer graded final examination.

## Lung, colon, breast and prostate cancer account for half of the U.S. cancer mortalities.

| TYPE          | INCIDENCE | (MORTALITY) |
|---------------|-----------|-------------|
| Lung          | 171,900   | (157,200)   |
| Colon/Rectum  | 147,500   | (57,100)    |
| Breast        | 211,300   | (39,800)    |
| Prostate      | 220,900   | (28,900)    |
| <u>Others</u> | 582,500   | (273,500)   |
| Total         | 1,334,100 | (556,500)   |

Jemal, Ward and Thun, "Cancer: Principles & Practice of Oncology." Edited by DeVita, Hellman and Rosenberg. (2006), pp. 226-241

### Cancers which kill 10,000-30,000 U.S. patients annually include:

- Pancreatic cancer
- Non-Hodgkin's Lymphoma
- Leukemia
- Stomach cancer
- Ovarian cancer
- Brain cancer
- Liver cancer
- Bladder cancer
- Esophageal cancer
- Kidney cancer

### Cancer risks include:

- Alcohol
- Asbestos
- Diet
- Familial
- Hormones

### Cancer risks (continued)

- Obesity
- Ion Radiation
- Tobacco
- U.V. Radiation
- Viral

### Lung Cancer kills over 150,000 patients in the U.S. annually.

- There are 45 Million current smokers and 45 Million ex-smokers in the U.S.
- It is difficult to quit smoking due to nicotine addiction.

### Carcinogens which have been identified in cigarette smoke include:

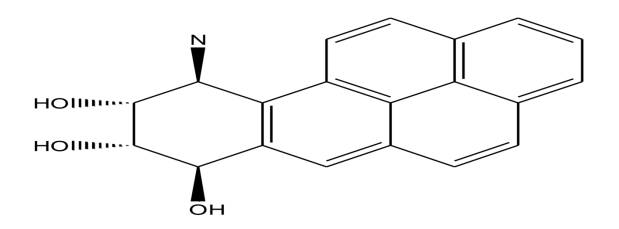
- Polyaeromatic hydrocarbons (PAH),
- aza-arenes,
- 4(methylnitrosamino)-1-(3-pyridyl)-1butanone (NNK),
- 1,3 butadiene,
- ethyl carbamate,
- ethylene oxide,
- nickel, chromium, cadmium,
- polonium, arsenic
- hydrazine

The process by which unreactive carcinogen converts to a form which binds DNA is known as metabolic activation.

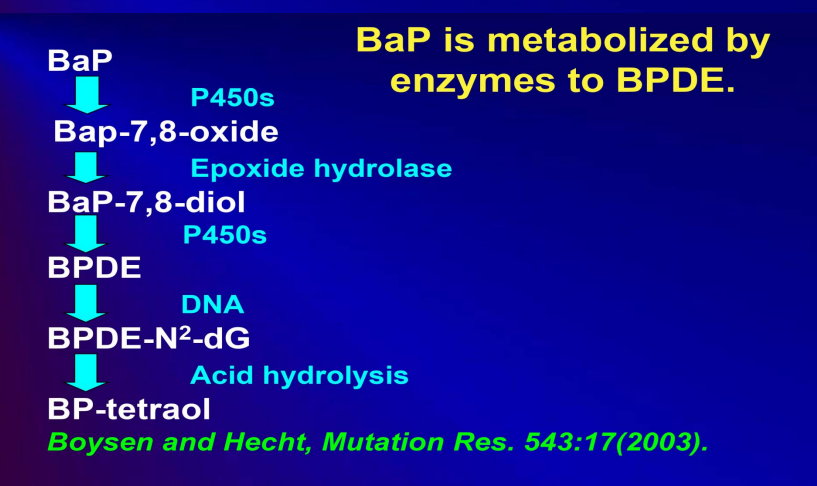
 Bay region diol epoxides are the principal PAH metabolites involved in DNA adduct formation. For Benz[a]pyrene (BaP), BaP-7,8-diol-9,10epoxide (BPDE) forms adducts with DNA leading to G:C>T:A mutations in pulmonary DNA. The genes for p53 and k-ras are frequently mutated.

### BENZ(a)Pyrene

### BENZ(a)Pyrene The chemical structure of BaP is shown.



### **BaP** is metabolized to BPDE



### Carcinogens can be detoxified and excreted prior to DNA damage.

- Cytochrome p450 enzymes catalyze addition of an oxygen to the carcinogen, increasing its water solubility.
- Phase 2 enzymes convert the oxygenated carcinogen to a form that is highly soluble in water, converting it to a form that can be excreted.

# DNA is mutated if the rate of carcinogen activation exceeds the rate of carcinogen detoxification and/or DNA repair.

 DNA adducts as well as intra- and interstrand DNA crosslinks are removed by nucleotide excision repair.

### P53, a tumor suppressor gene:

- mediates the G1 to S-phase checkpoint of the cell cycle,
- drives programmed cell death or apoptosis after DNA damage,
- is increased along with p21 (cell cycle checkpoint) after DNA damage.
- Phosphorylated p53 induces expression of BAX (apoptosis), GADD45 (DNA repair) and thrombospondin (angiogenesis)

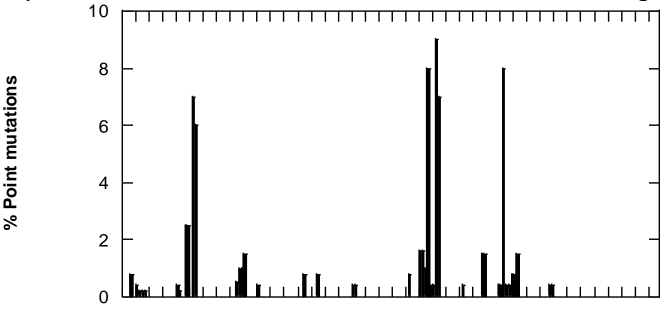
### P53 mutations are detected in most of the lung cancer patients.

 G to T transversions occur at the CpG rich codons including 153-158 (exon 5), 248 and 249 (exon7) and 273 (exon 8) of the p53 gene. There is an excess of G to T transversions in smokers relative to non-smokers.

#### P53 mutations.

P 53is mutated at codons 157, 158,
245, 248, 249 and 273 in lung cancer.

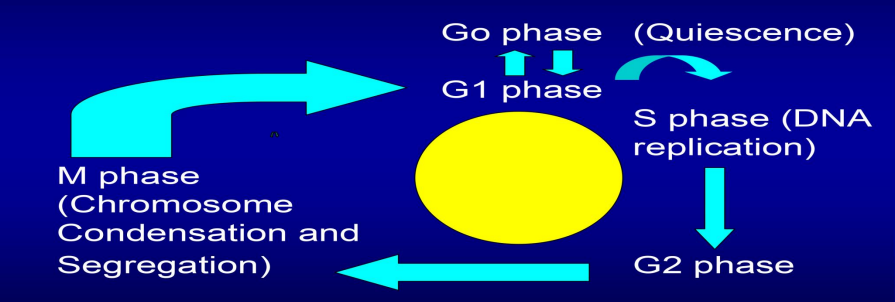




### Cell cycle phases

#### Cell cycle phases.

Cell cycle phases include G1, S, G2 and M



### p53 mediates the G<sub>1</sub> to S-phase checkpoint of the cell cycle

DNA damage increases p21 and p53.

 P53 drives programmed cell death or apoptosis after DNA damage

### Cell cycle enzymes

#### Cell cycle enzymes.

 Cyclin D/cdk is inhibited by p21,27,57,15,16,18 and 19.

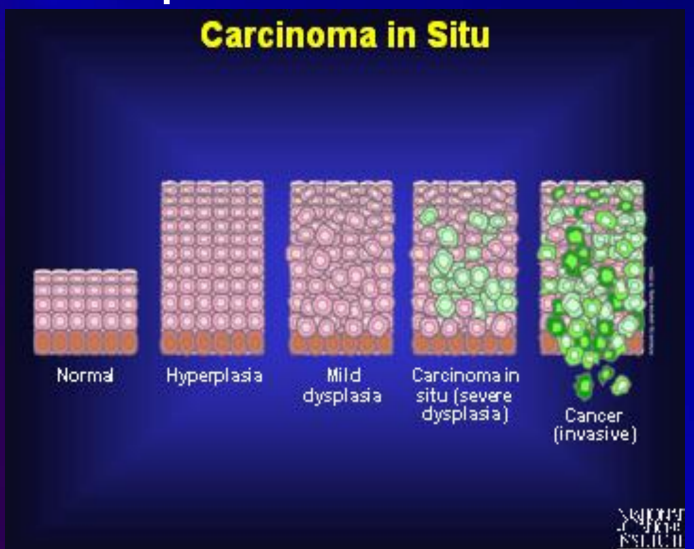


### Genotoxicity of tobacco smoke.

- After 10 years of chronic cigarette smoking, normal lung tissue can undergo hyperplasia and metaplasia.
- After 15 years, dysplasia can result.
- After 20 years, a carcinoma in situ can form.
- After 25 years, a malignant cancer can form.

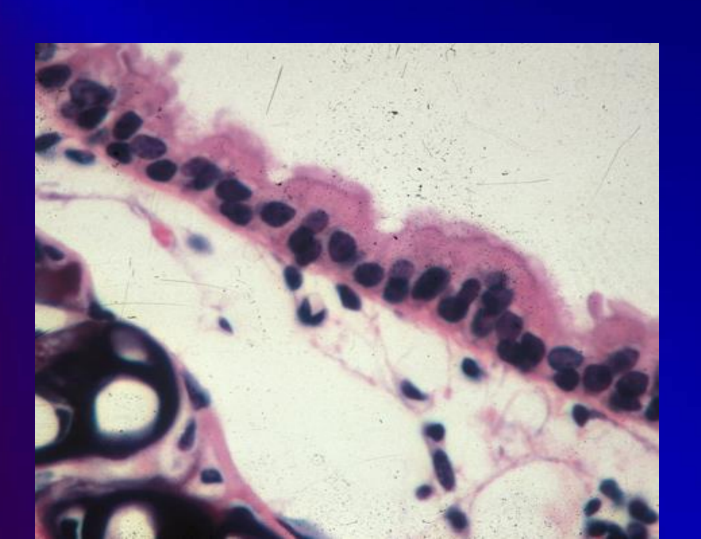
### Carcinogenesis

Cancer progression occurs over a period of decades.



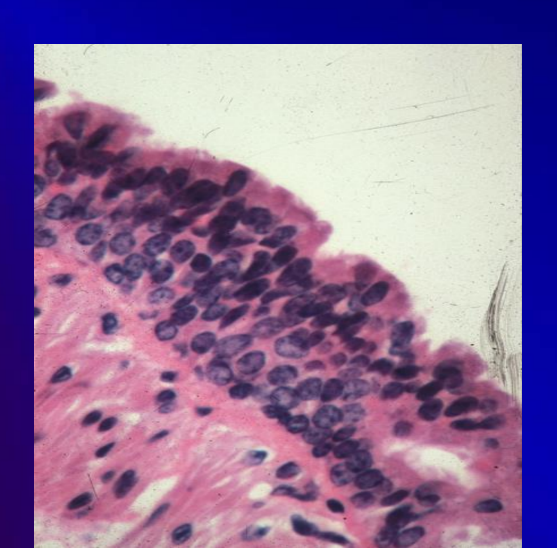
#### Normal lung

 Carbon dioxide is exhaled from the lung whereas oxygen is inhaled.



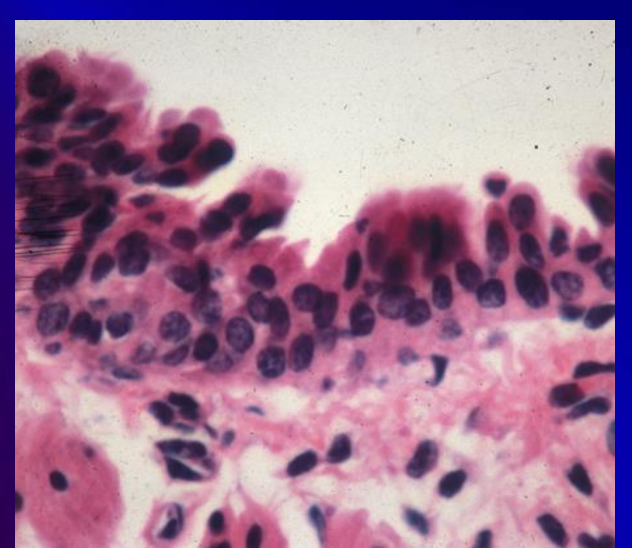
### Hyperplasia

• After exposure to tobacco smoke, hyperplasia can result.



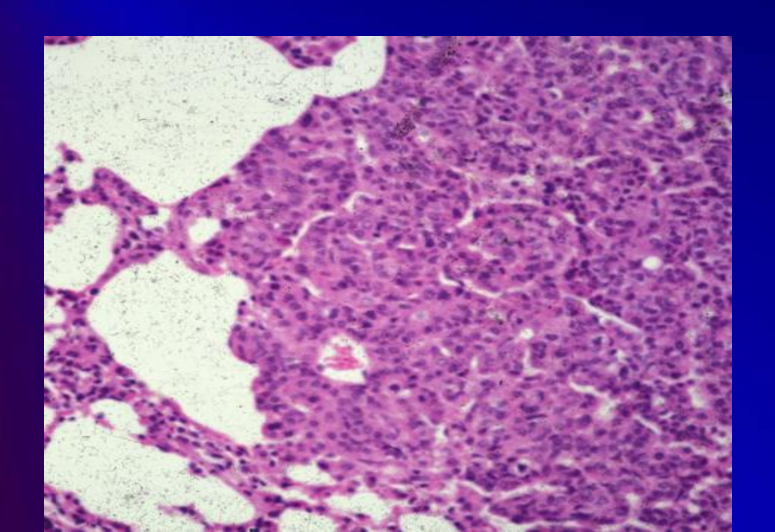
### **Dysplasia**

Continued exposure to tobacco smoke leads to dysplasia.



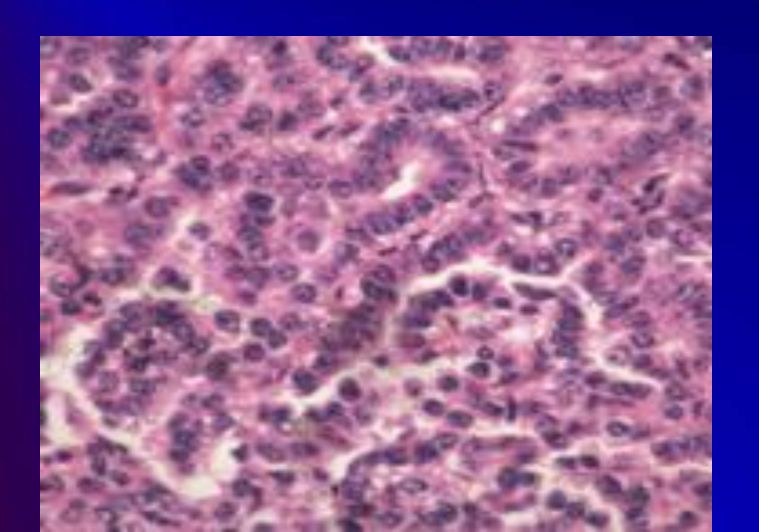
### **Adenoma**

 Continued exposure to carcinogens leads to benign tumors such as adenomas.



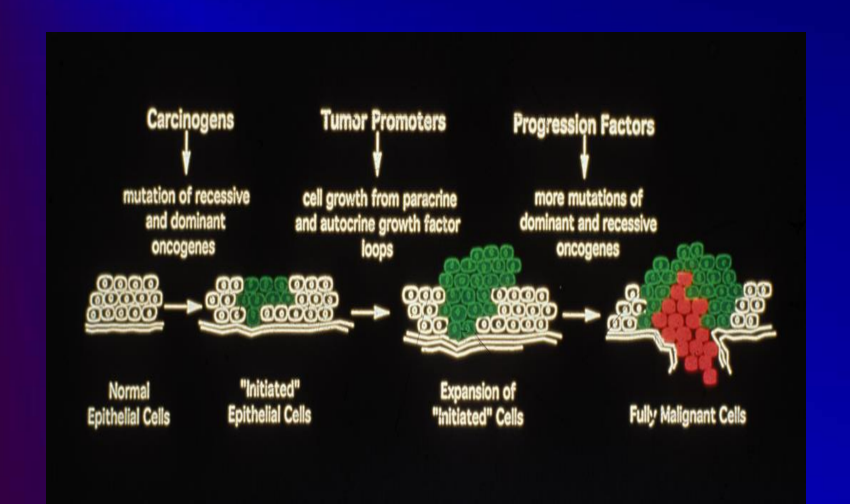
#### Adenocarcinoma

 Chronic exposure to tobacco leads to malignant tumors such as adenocarcinoma.



#### **Tumor formation**

Growth factors promote carcinogenesis.
 Progression factors lead to malignant tumors.



### Tumor growth

#### **Tumors**

 The primary cancer can undergo metastasis to distant organs.
 Carcinoma

**Angiogenesis** 

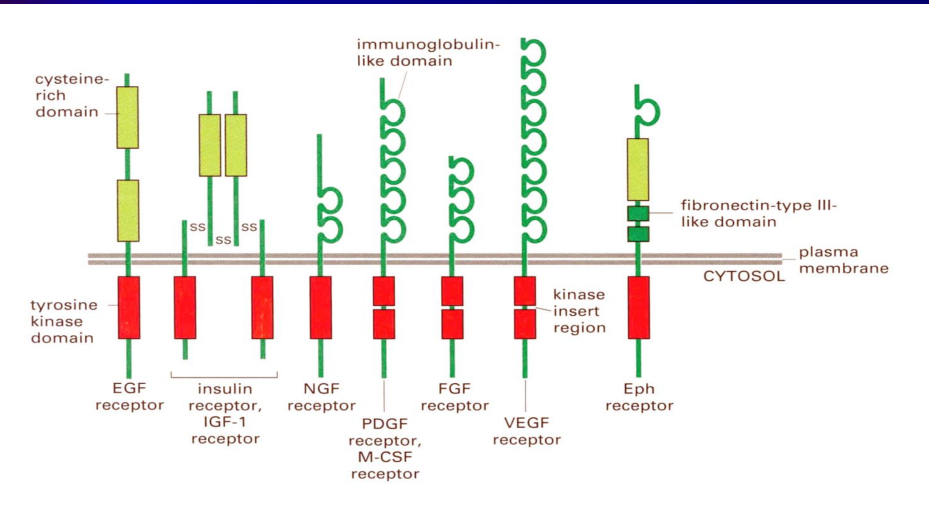
Migration, Invasion and Metastasis.

### Genetic abnormalities in lung cancer include:

- Mutation of tumor suppressor genes such as p53
- Silencing of tumor suppressor genes such as p16, Rb
- Amplification of oncogenes such as c-myc, cyclin D1, EGF receptor, erbB-2

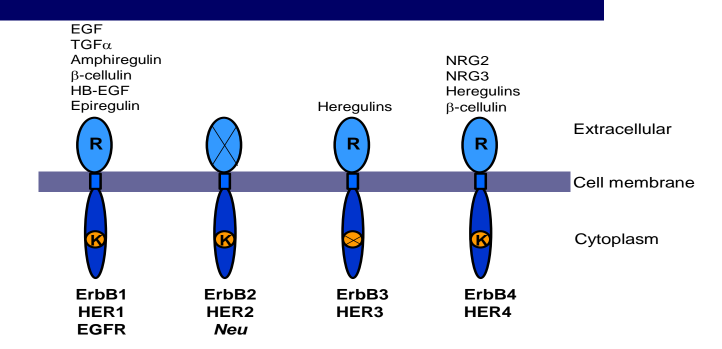
## Tyrosine kinase receptors.

Molecular Biology of the Cell, Alberts et al., 2001.



## Tyrosine kinase receptors and ligands

## **ErbB** family of receptor tyrosine kinases (RTKs) and ligands



## The EGFR is an 1186 amino acid integral membrane protein.

- The 621 amino acid extracellular domain binds EGF with high affinity. Domains I and III form the EGF binding site whereas domains II and IV are enriched in cysteine amino acids.
- The 24 amino acid transmembrane domain anchors the receptor into the membrane and tranduces signaling.
- The 541 amino acid intracellular domain contains tyrosine kinase activity.
- Lys721 binds ATP and Tyr amino acids are subsequently phosphorylated.
- Tyr1068, 1086, 1148, 1174 are autophosphorylated

## EGF, TGFα and mAb 108 bind with high affinity to lung cancer cells.

Agent IC<sub>50</sub>, ug/ml

EGF .03

 $\mathsf{TGF}\alpha$  .8

TGF $\alpha$ -PE38 .4

mAb 108 3

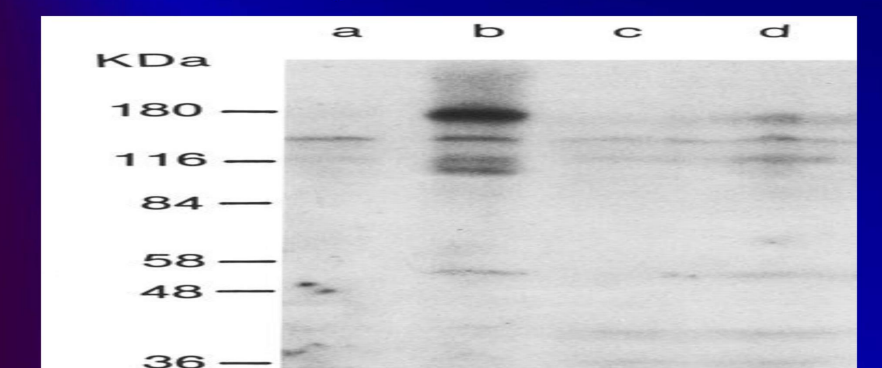
IgG >10

The IC<sub>50</sub> to inhibit <sup>125</sup>I-EGF specific binding to NCI-H157 cells was determined.

Draoui et al., Life Sci. 1994; 35:352.

## EGF tyrosine phosphorylation

EGF causes tyrosine phosphorylation of the EGFR, PLC<sub>γ</sub>, and PI-3-K.

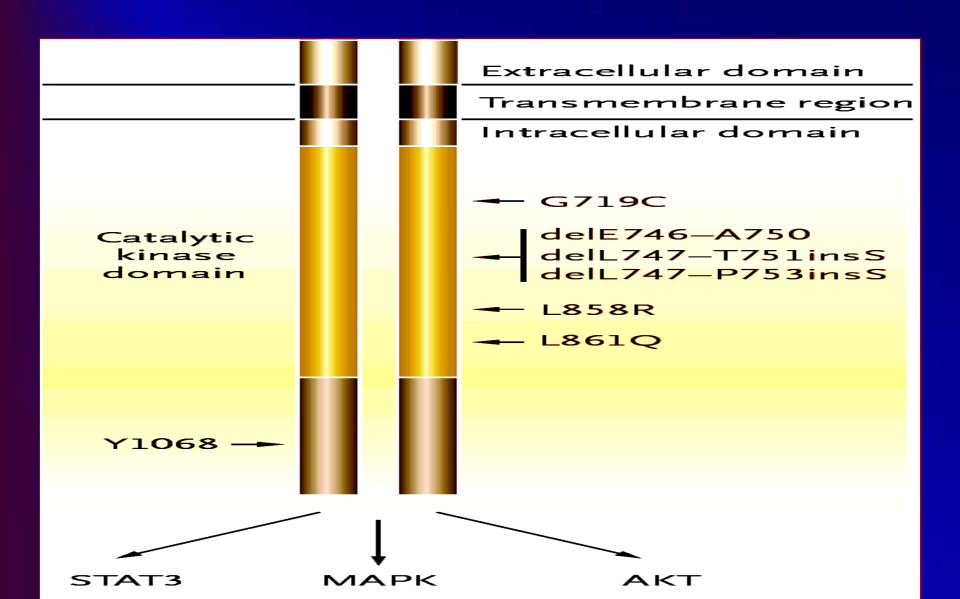


# Tyrosine kinase receptors are mutated in several diseases leading to increased cancer proliferation.

- EGFR mutations occur in the activation loop, especially L858R and G719C.
- Tyrosine kinase inhibitors (gefitinib and erlotinib) have been developed for the EGFR.

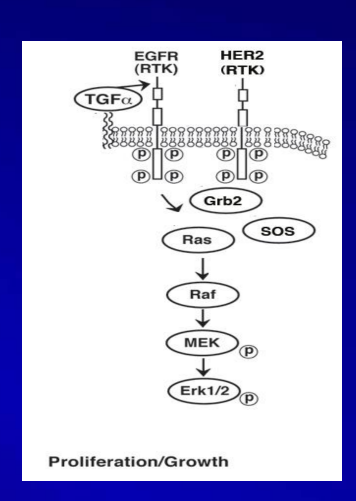
Paez et al., Science 304:1497 (2004)

## **EGFR** mutations



## RAS, RAF, MEK and ERK

- Receptor tyrosine kinases (RTK) stimulate proliferation
   Through the RAS, RAF,
   MEK and ERK pathway
- In NSCLC, K-RAS is
   Mutated in approximately 20%
   of the patients.



### RAS

- Mutated RAS has reduced GTPase activity resulting in an abundance of biologically active RAS-GTP.
- Most of the RAS mutations are G-to-T transversions in codon 12.
- The Frederick National Lab has a new initiative with RAS as a molecular target.

### **RAF**

- RAF is a serine threonine kinase which activates MEK. B-RAF-V600E mutations occur in approximately 60% of melanoma patients leading to an active kinase.
- PLX4032 is a kinase inhibitor which has an 81% response rate in patients with metastatic melanoma.
- RAS and B-RAF are driver mutations in several types of cancer

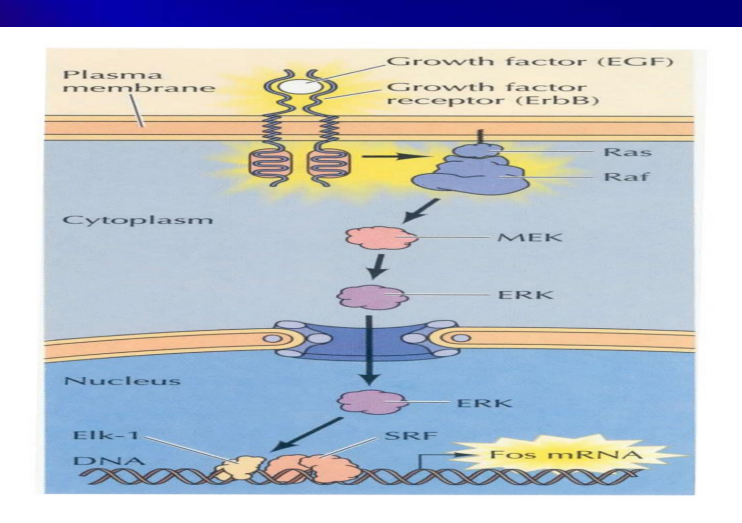
### MEK

- RAF phosphorylates mitogen activated protein kinase kinase (MEK) increasing its activity.
- MEK1 and MEK2 are inhibited by trametinib in B-RAF inhibitor –naïve patients.
- The MEK1/MEK2 inhibitor selumetinib plus docetaxel are being investigated in KRAS-mutant NSCLC patients.

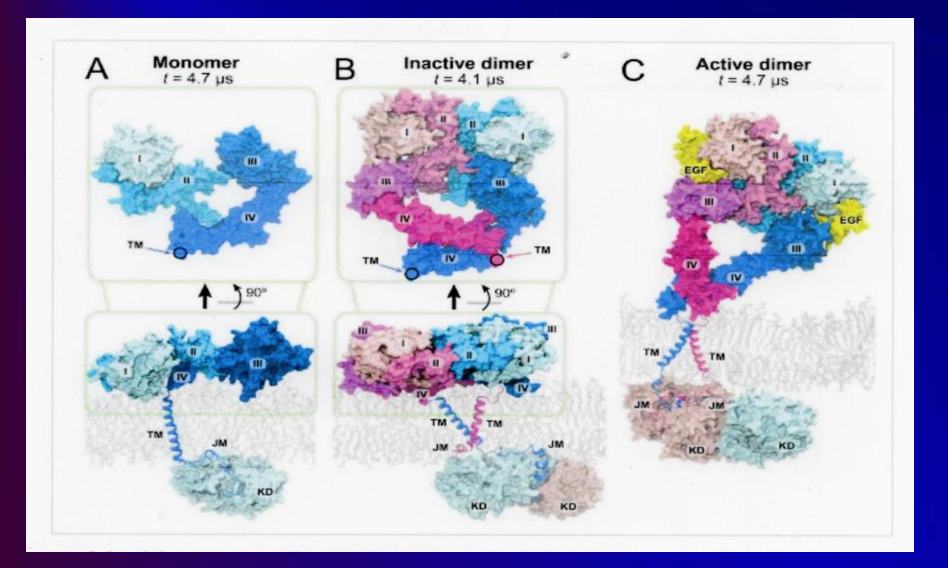
## **ERK**

- •MEK1/MEK2 regulates the phosphorylation of extracellular signal-regulated kinases (ERK) 1 and 2.
- •Phosphorylated ERK goes to the nucleus where it regulates expression of transcription factors such as fos, jun or myc.

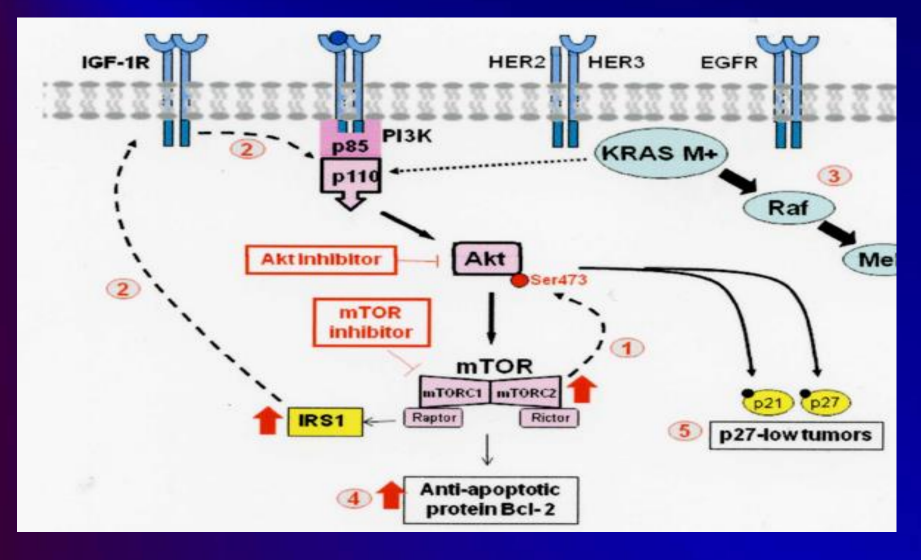
## The EGFR stimulates cancer cell growth. Molecular Biology of the cell; Alberts et al., 2001.



## **EGFR** dimerization



## PI3K, Akt, mTOR pathways stimulate cellular survival.



### PI3K

- The phosphatidylinositol 3 kinase (PI3K) pathway promotes cancer cell survival.
- The catalytic 100 kDa subunit metabolizes PIP<sub>2</sub> to PIP<sub>3</sub>
- PI3K is mutated in breast (25%), brain (27%), colon (30%) and stomach (25%) at E542, E545 or H1047 resulting in a gain of enzymatic activity.

### **PTEN**

- PI3K mutations involve chromosome 10q, which contains phosphatase and tensin homolog (PTEN).
- PTEN metabolizes PIP<sub>3</sub> to PIP<sub>2</sub> leading to inhibition of AKT signaling.
- PTEN is mutated in approximately 13% of breast cancer patients but loss of heterozygosity is more common.

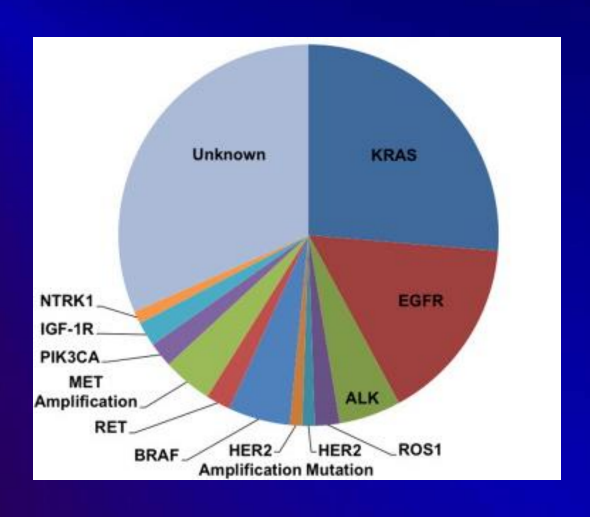
### Akt

- AKT or protein kinase B prevents apoptosis of cells.
- AKT is a serine/threonine kinase which is phosphorylated at Ser473 increasing phosphorylation of mTOR.
- AKT promotes cellular survival by phosphorylating BAD and caspase-9 preventing apoptosis of cancer cells.
- AKT is mutated in breast cancer (5%), colorectal cancer (6%) and ovarian cancer 2%.

### **mTOR**

- Mammalian target of rapamycin (mTOR) or FRAP1 is a serine/threonine kinase.
- mTOR activation enhances phosphorylation of p70S6 kinase and 4E-BP1 increasing protein translation and cellular proliferation.
- mTOR activation decreased autophagy, a lysosome-dependent degradation pathway.

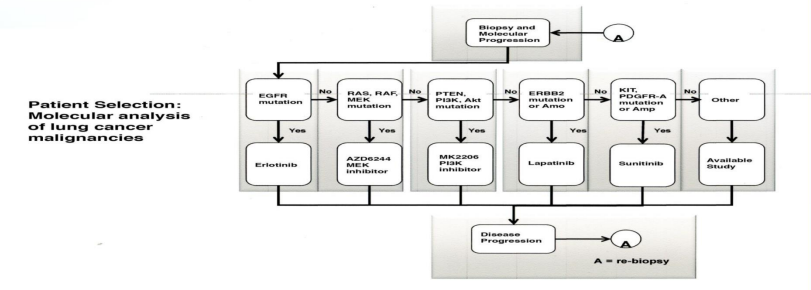
## Personalizing Therapy for NSCLC Genetic Abnormalities in Lung Adenocarcinoma



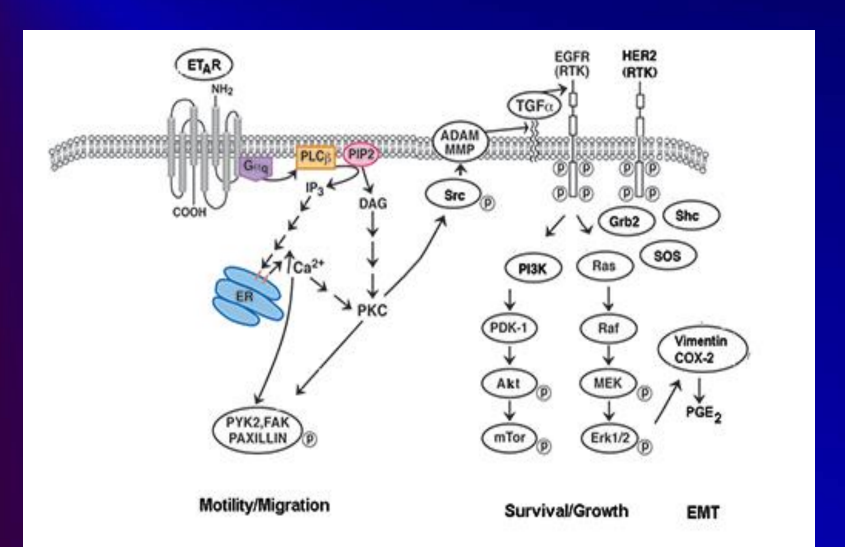
## Molecular medicine

## Molecularly Targeted Treatment of Advanced Thoracic Malignancies

#### Molecularly Targeted Treatment of Advanced Thoracic Malignancies



## GPCR transactivate the EGFR and HER2



## Erlotinib/gefitinib resistance

 Approximately 50% of NSCLC patients develop resistance to erlotinib/gefitinib after 1 year due to a secondary mutation in the EGFR (T790M).

## CML patients are sensitive to the small molecule TKI Gleevec.

 This restores blood counts in patients and delays disease progression.

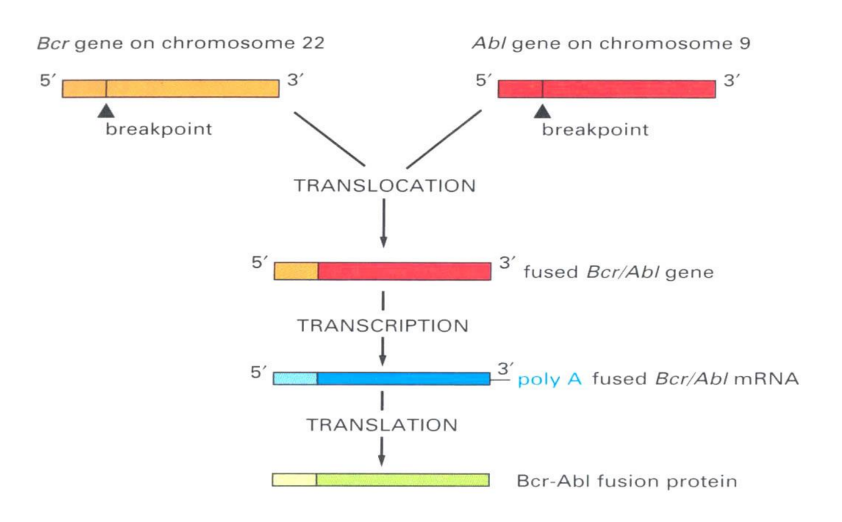
## CML patients

## CML patients have a genetic abnormality on chromosome 22 (Philadelphia chromosome).

- Segments of chromosome 9 and 22 are fused resulting in the bcr-abl gene. ●The resulting tyrosine kinase is constituitively active.
- Bcr-abl tyrosine kinase actvity is inhibited by Gleevec.

#### Translocation of Bcr/Abl.

• Chromosome 22 translocates with chromosome 9. Molecular Biology of the Cell; Alberts et al., 2001.

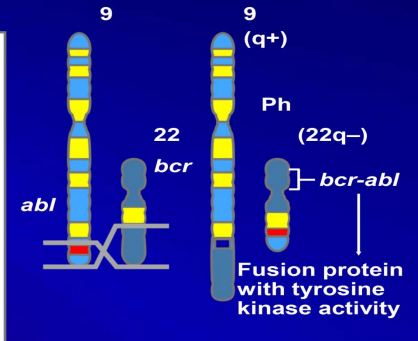


### Bcr-Abl

#### Translocation of *Bcr-Abl* Genes

 Translocated chromosome 9 appears larger and translocated chromosome 22 appears smaller: Freebies for Teachers"; D. Kerrigan.





Artwork by Jeanne Kelly. © 2004.

# In a Phase I Clinical Trial, Gleevec<sup>TM</sup> was effective orally at a daily dose of 300 mg or greater.

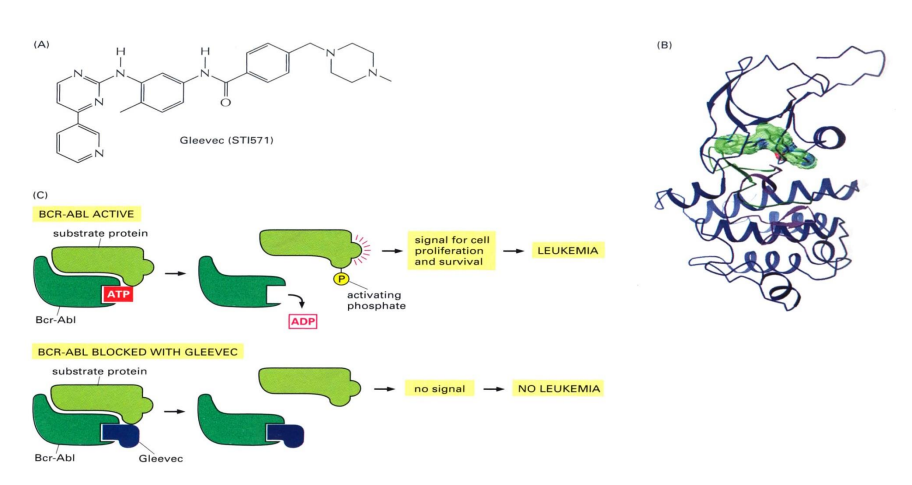
 Dose limiting toxicities included nausea, vomiting, edema and rash. (Sawyers and Druker. Cancer J. Sci. Am. 1999;5:63).

# In a Phase II Clinical Trial, Gleevec<sup>TM</sup> restored normal blood counts in 53 out of 54 chemotherapyresistant CML patients.

- After a year on Gleevec, 51 of these patients were still doing well. (Druker et al. N. Engl. J. Med. 2001; 344: 1038.).
- Over a 5 year period, 89% of the patients treated with Gleevec had progression-free survival (O'Hare et al., Clin. Cancer Res. 2011; 17: 212).

#### Gleevec mechanism of action

•Gleevec blocks the ATP binding site. Molecular biology of the cell; Alberts et al., 2001.



#### **GLEEVEC RESISTANCE**

- •Over a 5 year period, 17% of the patients initially sensitive to Gleevec became resistant.
- BCR-ABL point mutations occurred such as T315I near the ATP binding site impairing Gleevec interactions
- New drugs such as ponatinib or DCC-2036 are being developed which bind with high affinity to mutated BCR-ABL

## Tyrosine kinase inhibitors in cancer

CML Bcr-Abl Imitanib/dasatanib

Breast cancer HER2 Herceptin/lapatanib

Melanoma B-RAF PLX4032

GIST c-KIT Imatinib/sunitinib

NSCLC EGFR Gefitinib/erlotinib

## PRACTICAL STEPS TO PREVENT CANCER

- Check your house for radon.
- Check your house for asbestos.
- Take precautions at your workplace.
- Check your community water system.
- Avoid breathing polluted air.
- Protect your skin.
- Don't breathe smoke.
- Exercise daily.

## **Cancer Prevention**

## PRACTICAL STEPS TO PREVENT CANCER (continued)

- Avoid pesticides.
- Eat fruits and vegetables.
- Reduce red-meat consumption.
- Eat fish.
- Minimize fried foods.
- Drink alcohol in moderation.
- Avoid unnecessary x-rays.
- Reduce infections.

### REFERENCES

#### REFERENCES

- Hanahan, D. and Weinberg, R.A. Hallmarks of cancer: The next generation. Cell 2011; 144(5): 646-74.
- ●O'Hare, T., Deininger, M.W.N., Elde, C.A., Clackson, T., and Druker, B.J. Targeting the BCR-ABL signaling pathway in therapyresistant Philadelphia chromosome-positive leukemia. Clin. Cancer Res. 2011; 17(2):212-21.