GENENTECH INC Form DEFA14A November 17, 2006

UNITED STATES

SECURITIES AND EXCHANGE COMMISSION

Washington, DC 20549

SCHEDULE 14A

Proxy Statement Pursuant to Section 14(a) of the

Securities Exchange Act of 1934

Filed by the Registrant x

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Check the appropriate box:

" Preliminary Proxy Statement

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Genentech, Inc.

(Name of Registrant as Specified in its Charter)

(Name of Person(s) Filing Proxy Statement, if other than the Registrant)

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x No fee required.

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#1Genentech: An OverviewGenentech: An OverviewPatrick Yang, Executive Vice PresidentManufacturing, GenentechNovember 17, 2006

#2 Meeting Agenda Introduction to Genentech Pat Yang Manufacturing Next steps Pat Yang Q&A All

#3 Manufacturing Manufacturing

Genentech is a world leader in biotech manufacturing, with more FDA-approved manufacturing capacity for the production of biotech medicines than any other company

Four facilities: South San Francisco, CA; Vacaville, CA and Oceanside, CA and Porriño, Spain

We believe we have the right plans in place to meet the growing demand for our products:

Oceanside facility purchased from Biogen Idec in 2005

Option to purchase facility in Singapore

Working with Lonza, Wyeth and Novartis

Process yield improvements for Rituxan and Avastin

New capacity coming online for bulk and filling/packaging

#4

Q4 06, announced Lonza will acquire our 40,000 liter facility in Porriño, Spain; Lonza will continue production of Avastin for Genentech for 3 years

We anticipate closing the transaction before the end of 2006 Porriño, Spain (Lonza Biologics) Status **Enhancement Project** Facility Anticipate construction, qualification and licensure of our new plant in Vacaville, California in 2H 09 (additional 200,000 liters) Vacaville, CA CCP2 Genentech Bulk Manufacturing Oceanside, CA NIMO Anticipate FDA licensure to produce commercial Avastin in 1H 07 (90,000 liters) Contract Manufacturing Wyeth BioPharma Andover, MA Received FDA licensure to produce Herceptin Q3 06 Process Improvements Rituxan Anticipate approval of higher titer Rituxan process in Vacaville by the end of 2006 (+50%)Avastin Anticipate approval of higher titer Avastin process in South San Francisco by the end of 2006 (+50%) Near-term Key Capacity Enhancement Projects Near-term Key Capacity Enhancement Projects As of November 9, 2006

#5 Novartis Pharmaceuticals Huningue, France

Began manufacturing all future worldwide supply of Xolair Wyeth BioPhmara Andover, MA

Received FDA licensure to produce

Herceptin Lonza Biologics Porrino, Spain

Announced Lonza will acquire our 40,000 liter facility in Porriño, Spain; facility will continue production of Avastin for 3 years Lonza Biologics Singapore

Entered into long-term supply agreement with Lonza to manufacture Genentech products at their 80,000-liter facility in Singapore; We have an exclusive option to purchase the Singapore facility in the future 2006 Key Contract Manufacturing Accomplishments 2006 Key Contract Manufacturing Accomplishments

Our strategies include expanding or acquiring facilities and engaging contract manufacturers that produce Genentech s products on our behalf As of November 9, 2006 #6

Completed qualification runs of Avastin at Oceanside

Expect FDA licensure to produce Avastin 1H 07

Purchased state-of-the-art finish/fill facility in Hillsboro, Oregon

Expect facility to be licensed and operational in 2010

Expect FDA approval of high titer processes for Rituxan (in Vacaville) and Avastin (in SSF)

Signed two new product supply agreements with Roche

Other 2006 Manufacturing Accomplishments Other 2006 Manufacturing Accomplishments As of November 9, 2006

#7

Genentech s Oceanside Facilities Genentech s Oceanside Facilities Manufacturing Facility Manufacturing Facility NIMO **Commercial Facility** NICO **Clinical Facility** Purchased Purchased June 2005 February 2006 Potential Capacity Potential Capacity 90,000 liters 5,500 liters # of Employees # of Employees Approximately 530 employees as of September 30, 2006 Plan to employ approximately 30 employees by the end of 2006 Status Status

Q3 06 completed qualification runs of Avastin

Expect FDA licensure to produce Avastin in 1H 07

Expect to be operational by Q1 07 QuickTime and a MPEG-4 Video decompressor are needed to see this picture.

#8 -

Potential to purchase Lonza Singapore Facility 610,000 Liters Option to Purchase from 2007

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2010 Q4 06Genentech obtained an exclusive option to purchase the Lonza Singapore facility during the period from 2007 to 2012 Licensure to produce Avastin is expected 2010 80,000 liters Lonza Singapore Facility Q4 06. announced Lonza will acquire our 40,000 liter facility in Porriño, Spain; Lonza will continue production of Avastin for Genentech for 3 years. We anticipate closing the transaction before the end of 2006. Lonza Biologics, Porriño, Spain Comments Other Comments **Potential Capacity** Genentech Bulk Manufacturing Facility 240,000 Liters Current Total Capacity in Use Potential addition of Vacaville, CA 530,000 Liters Potential Capacity in 1H 09 Potential addition of Oceanside, CA 330,000 Liters Potential Capacity in 1H 07 Comments Contract Manufacturing (Bulk) Expect FDA licensure in 2H 09 200,000 (8x25,000L) Vacaville, CA (CCP2) In July 2006, Roche signed two new product supply agreements which supplement and supersede existing produ agreements.

Roche
has
agreed
to
purchase
specified
amounts
of
Herceptin,
Avastin
and
Rituxan
through
2008
and
to
purchase
specified
amounts
of
Herceptin
and
Avastin
through
2012.
Previously Roche had assumed most of their own ex-US Hercentin supply and was planning on assuming all their ex-

Previously, Roche had assumed most of their own ex-US Herceptin supply and was planning on assuming all their ex-US Avastin supply.

Genentechas and will continue to supply all of Roche s ex-U.S. Rituxan supply.

Roche, Penzberg, Germany

Received FDA licensure in Q1 06 to produce bulk substance olair (will produce all future worldwide supply). As o Genentech will

acquire bulk supply of Xolair from Novartis and compensate them on a cost plus mark

up basis. Novartis Pharmaceuticals, Huningue, France Received FDA licensure in Q3 06 to produce Herceptin; expect Wyeth to produce 25% of Herceptin over the next several years. Genentech will produce the remainder in Vacaville. Wyeth BioPharma, Andover, MA Received FDA licensure in Q3 05 to produce Rituxan; expect Lonza to produce ~50% of Rituxan over the next set Genentech will produce the remainder in our other facilities. Q3 06 we completed qualification runs of Avastin Expect **FDA** licensure to produce Avastin in 1H 07 First licensed in 2000. Licensed to produce Avastin, HerceptinRituxan, Xolair. First licensed in 1985. Licensed to produceActivase, Avastin, Cathflo Activase, Herceptin, Lucentis, Nutropin, Nutropin AQ, Pulmozyme, Raptiva, Rituxan, and TNKase. Comments Lonza Biologics, Portsmouth, NH 90,000 (6x15,000L) Oceanside, CA (NIMO) 144,000 (12x12,000L) Vacaville, CA (CCP1) 96,000 (8x12,000L)

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South San Francisco, CA Current Capacity Genentech Bulk Manufacturing Facility Manufacturing Capacity Manufacturing Capacity As of November 9, 2006

#9 0 50,000 100,000 150,000 200,000 250,000 300,000 350,000 400,000 450,000 500,000 550,000 600,000 650,000 1999

2000 2001 2002 2003 2004 2005 2006 1H'07 2008 2H'09 2010 *Chinese Hamster Ovary Cell Culture Note: In Q4 06, Genentech has entered into an agreement with Lonza to purchase Genentech's Porrino, Spain manufacturing facility. Concurrently, we entered into a supply agreement for the manufacture of certain Genentech products at Lonza's facility currently under construction

in Singapore, with Genentech also receiving the right to exercise an exclusive option to purchase the Lonza Singapore facility during the period from 2007 to 2012. The transactions are subject to various closing conditions.. As of November 9, 2006 Genentech Commercial Cell Culture* **Bioreactor Capacity** Genentech Commercial Cell Culture* **Bioreactor Capacity Current Commercial Capacity** South San Francisco, CA and Vacaville, CA (CCP1) #10 What Does This Mean For You? What Does This Mean For You?

We encourage continued focus on your current efforts to bring important new medicines to patients, as this is in everyone s best interest

While we fully expect the deal to go through, we aren t there yet

Your current management continues to run the company until close

Once the GNE and Tanox transition teams are up and running, more detailed information will be available on next steps, key milestones, etc.

Most importantly, we recognize that this is an uncertain time for Tanox employees. Consistent with our values, our intent is to treat Tanox employees with the same respect & integrity that we treat our own employees

#11
Transition Process Will Be Organized Around Four Areas
Transition Process Will Be Organized Around Four Areas
EC / Legal
EC
Product
Portfolio
Committee

Research Review Committee EC / PROP **Executive Team** Decision maker Feb. 28, 2007* Feb. 28, 2007* Jan. 31, 2007* Feb 28, 2007* 3.Recomm end-ation Feb 10, 2007* Feb. 10, 2007* Jan. 31, 2007* Jan 31, 2007* 2. Evaluation Number of contracts; Rights and obligations of each Number of employees by functional area Number and value of R&D programs Number, location, and capability of facility 1. Assessmen t Contracts HR Change management R&D Programs Facilities/ Property **Process Summary** * All dates are tentative #12 Decisions Yet to be Determined Decisions Yet to be Determined

Future plans for Tanox s pipeline

Future plans for Tanox s sites

Future status of employees

Who will be retained

How/when decisions will be made after close; however, our intent is for decisions to be made as quickly and with as much transparency as possible

Where retained employees will be located

Details of post-close integration and timeline

#13 Next Steps Next Steps Today:

Small functional meetings with Genentech and Tanox management Next few months: