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## Optimization of the Oxidant Supply System for Combined Cycle MHD Power Plants

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Work performed for U.S. DEPARTMENT OF ENERGY Fossil Energy Office of Magnetohydrodynamics

Prepared for Twentieth Symposium on the Engineering Aspects of Magnetohydrodynamics Irvine, California, June 14-16, 1982

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# Optimization of the Oxidant Supply System for Combined Cycle MHD Power Plants

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Work performed for U.S. DEPARTMENT OF ENERGY Fossil Energy Office of Magnetohydrodynamics Washington, D.C. 20545 Under Interagency Agreement DE-AI01-77ET10769

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#### OPTIMIZATION OF THE OXIDANT SUPPLY SYSTEM FOR COMBINED CYCLE MHD POWER PLANTS

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#### Abstract

An in-depth study was conducted, both under contract and in-house, to determine what, if any, improvements could be made on the oxidant supply system for combined cycle MHD power plants which would be reflected in higher thermal efficiency and a reduction in the cost of electricity, COE. The study included a systematic analysis of "airseparation" process variations which showed that the specific energy consumption could be minimized when the product stream oxygen concentration is about 70 mole percent. The use of advanced air compressors, having variable speed and guide vane position control, results in additional power savings. The study also led to the conceptual design of a new air separation process, sized for a 500 MW<sub>e</sub> MHD plant, referred to as "internal compression." In addition to its lower overall energy consumption, potential capital cost savings have been identified for air separation plants using this process when constructed in a single large air separation train rather than multiple parallel trains, typical of conventional practice.

The effect of the lower energy consumption of the new air separation process on major MHD plant parameters such as thermal efficiency and magnetic energy requirement was calculated for a range of channel lengths and for various oxidant stream compositions. The process was shown to bring about an increase in overall plant thermal efficiency of 0.6 percent while allowing more favorable tradeoffs between magnetic energy and oxygen system capacity.

#### Introduction

Oxygen enrichment of the MHD combustor oxidant stream at low and intermediate preheat temperatures has been identified as a viable alternative to systems using indirectly fired high temperature air heaters. 1, 2, 3 These studies identified the attractiveness of MHD/steam power plants using intermediate temperature recuperative preheat and oxygen enrichment of the combustor oxidizer for early commercial MHD power plants. Detailed MHD plant performance calculations considering oxygen enriched MHD combustion air4 showed that the energy consumption required for oxygen production has a significant impact on the overall results. Although energy consumption values required for oxygen production by air separation plants delivering a high purity oxygen product (above 99 mol percent 02) range from 280 to 300 kWh/ton of pure 02, considerable energy savings can be achieved by using a medium purity product (60 to 80 mol percent 02) for blending with air to produce the required MHD combustor oxidant stream whose oxygen concentration may range from 25 to 40 mol percent.

Such a "medium purity" air separation plant was placed in service by Linde A.G. for the Thyssen Steel Works, Germany, in 1973.5,6 This plant delivers a 70,000 normal cubic meters per hour (ncmh) (43,500 scfm) product stream at near atmospheric pressure containing 60 mol percent oxygen which is blended with air to form a blast furnace stream of 283 000 ncmh (176,000 scfm) containing 30.6 mol percent oxygen. The plant's specific energy consumption (SEC) is reported at 224 kWh per ton of equivalent pure oxygen (TEPO).

The present study was managed by NASA LeRC for DOE under Interagency Agreement No. EF-77-A-01-2674. The Lotepro Co. of New York, NY a subsidiary of Linde A.G., FRG (Germany) was the main contractor under contract DEN3-165. Also supplying data were Linde A.G. and a number of industrial compressor manufacturers; notably, Demag-Mannesman Co., Dresser Industries, Inc., MAN/GHH Co., Sulzer Brothers, Inc., and Transamerica De-Laval, Inc.

The study objectives and preliminary results have been reported on previously in its early stage. The main objective was to explore process variations for their potential reduction in SEC below the 224 kWh/TEPO quoted above. The work has recently been completed, and a comprehensive report is in preparation. The present paper summarizes the main results of the study, including a brief description of a new air separation process applicable to thermodynamic cycles requiring pressurized oxidant streams, and its effect on the performance of a 500 MWe combined cycle MHD power plant.

#### II. Cryogenic Air Separation Process

Before discussing the study results, a brief review of the basic cryogenic air separation process is in order.

Cryogenic air separation processes utilize the difference in boiling points of the various air components (principally  $N_2$  and  $O_2$ ) and the changes in boiling points with pressure to achieve component separation by fractional distillation. The basic process utilizes a number of auxiliary components (compressors, expanders, heat exchangers, adsorbers, switching valves) whose function is to change the state of the input air stream so that component separation can take place, and also to condition the output streams (product oxygen and nitrogen) to user requirements. The actual separation or rectification process is usually carried out in "double rectification columns" consisting of an upper compartment operated at low pressure (near atmospheric in most cases) and a lower compartment operated at higher pressure. The lower compartment is usually referred to as the medium pressure column, the upper, as the low pressure

column. Both columns are equipped with a number of perforated .eparation trays, the use of which brings about a series of fractional evaporations and condensations between a rising vapor, that is successively enriched in nitrogen, and a falling liquid, which is being enriched in oxygen at each stage. The process requirement for this heat and mass transfer to occur in the two phase-two component countercurrent flow environment dictates the pressure difference between the two columns and thereby specifies the process energy input required for the Air Separation Unit (ASU) air compressor. This is because nitrogen, which at atmospheric pressure has a lower boiling point than the desired product oxygen, must condense at the top of the medium pressure column in order to evaporate product oxygen at the bottom of the low pressure column. The only way this requirement can be met is for the condensation of No to take place at a higher pressure than that of the evaporating product 02. It should be noted that the higher the desired 02 concentration (purity) of the product, which is collecting at the bottom of the low pressure column, the greater the pressure difference requirement between the two columns and the higher the compressor power input for the same input air flow.

#### Basic Air Separation Plant

A schematic of a conventional air separation plant based on the process described above is shown in figure 1. Its main components are:

Air compressor, (CP)

- Adsorbers for the removal of water and

carbon dioxide, (WS, MS1, MS2) - Expansion turbine for the production of

refrigeration, (T)

- Heat exchangers for cooling the air to liquefaction temperature and warming the oxygen and nitrogen products to ambient temperature, (E)

 Rectification columns, (C1 and C2) and condenser-evaporator, (K) for the separation of air into O2 and N2 on the basis of vapor-to-liquid heat exchange and mass

In many plants the removal of water and carbon dioxide is combined with the cooling of the air. This is done in reversing heat exchangers or regenerators. The flow paths of the air and product streams can be traced on figure 1. Filtered atmospheric air is compressed in an intercooled and aftercooled air compressor, CP, to a pressure which is determined by the requirements of the particular process, usually 4 to 6 bars. It is then passed through the water separator, WS, and adsorbers, MS, to the heat exchangers, E, and cooled to about  $100^{0}$ K with the aid of expansion turbine,  $\uparrow$ . The cold air is injected as a saturated vapor into the mediumpressure column, Cl, where it is separated into a nitrogen fraction and the so-called rich liquid fraction, which contains about 40 percent oxygen. The separation is obtained by contacting the rising vapor with liquid nitrogen flowing down from the condensor-evaporator, K. This causes the liquid collecting in the kettle at the bottom of C1 to become enriched in oxygen while the vapor, which is condensed at the top of Cl, is enriched in nitrogen. Both fractions leave Cl in

the liquid state and flow through expansion valves to the low-pressure column, C2, where additional rectification to the final product purity takes Roth nitrogen and product oxygen leave column C2 in the gaseous state at near atmospheric pressure and are heated in heat exchanger E before being ducted to the user. Note that for applications, such as MHD, requiring lower O2 concentrations than those of typical medium purity ASU products, the product stream is mixed with atmospheric air before compression in specially designed uncooled oxygen-enriched air compressors as indicated by oxidant compressor, OC, in figure Moreover, as shown in the appendix at the end of this paper, the mixing of air separation plant product and air, to obtain mixtures containing certain mole percentages of O2, will result in slightly different mixture compositions than those obtained by mixing pure oxygen with air.

#### III. Process Study and Results

#### Summary of Process Variations

The approach used was to do a systematic analysis of as many as 18 process variations for an oxidant supply system sized for a 1000 MW<sub>e</sub> MHD/steam power plant. The oxidant supply system included the air separation plant as well as the final oxidant mixing chamber and oxygen enriched air compressor. Delivery pressure of the oxidant stream to the MHD combustor was specified at 8 atm and the O<sub>2</sub> concentration of the oxidant stream was 30.6 mol percent. The process parameters varied in the study were product 02 concentration which was ranged from 30.6 to 99.5 mol percent 02, and the upper column pressure which was varied from 1 atm to 8 atm, with the high pressure cases also providing pressurized No for power recovery. For each process studied both total power consumption and relative cost figures were obtained.

Results showed that the atmospheric upper column pressure cases, in which the product is mixed with ambient air prior to compression to combustor pressure, were superior to the high pressure upper column cases, both in total power consumption and in capital cost. Figure 2 shows specific energy consumption, SEC, as a function of product purity for the atmospheric product cases. Note that the results represent a tradeoff between the ASU input air compressor pressure ratio requirement, which increases with product purity, and the ASU input air compressor flow requirement, which decreases with product purity. The SEC is shown to drop from 255 kWh/TEPO at 40 percent product purity to 195 kWh/TEPO at 70 to 80 percent purity and increase to near 240 kWh/TEPO for the high purity product. Hence the optimum (minimum compressor power) tradeoff between compressor flow and pressure ratio occurs at product purities between 70 to 80 percent. Since MHD applications require combustor oxidant 02 concentrations (25 to 40 mole percent) Well below the product purities at which compressor power is minimized, the more efficient medium purity air separation processes are compatible with MHD/steam power cycles. Based on these results a plant delivering a 70 mol percent O<sub>2</sub> product at one atmosphere was chosen for use in the 200 MW<sub>e</sub> Engineering Test Facility (ETF) Conceptual Design.<sup>8</sup>

#### New Air Separation Process

Having considered the available variations of the basic process exemplified by the Thyssen Steel plant, be decided to explore the potential additional improvement in total power consumption for a process which utilizes pumping of the product in the liquid state to the final end use pressure, product vaporization at this pressure, followed by heating and mixing with a separately compressed air stream to obtain the desired oxidant 02 concentration. Such a process may have significant advantages for end uses that require the process outout stream to be delivered at a pressure which is above atmospheric. The conceptual design for such a process was developed jointly by NASA LeRC and Lotepro. The process is referred to as an "Internal Compression and Liquid Product Pumping" in an invention disclosure currently being prepared.

#### Process Energy Consumption and Capital Cost

To evaluate the potential advantages of the new process over the previously discussed optimized conventional process a study was conducted of two oxidant supply systems, sized for a 500 MW<sub>e</sub> MHD/steam plant, using the new process and the optimized conventional process (195 kWh/TEPO), respectively. A comparison of power consumption results and of the main process parameters is shown in tables I and II. Also, a block diagram comparing the main features of the two oxidant supply systems is shown in figure 3.

Results in table I show that the new process, in which the product from the upper column is pumped as a liquid to the desired pressure, evaporated and heated, and then mixed with air compressed in the blend air compressor, has a lower overall energy consumption than the optimized conventional process in which an atmospheric pressure product is mixed with ambient air and the resulting oxidant stream is compressed in the oxidant compressor (see figure 3). Note that the oxidant compressor (an axial/radial machine) must be rated for oxygen enriched air service which implies a small penalty on efficiency and cost, whereas the blend air compressor used in the new process only has to compress air, which is less hazardous, and therefore can be compressed more efficiently in a multi-stage axial machine.

Even though, as is shown in table II, the SEC at product pressure for the new process is higher (267 vs. 195 kWh/TEPO), the overall energy required to produce the desired oxidant is lower. The use of the new process would be equivalent to using a conventional process, delivering product at atmospheric pressure, which has an SEC of 178 kWh/TEPO. This equivalent SEC is plotted as a triangle in figure 2. Further gains can be realized with the new process if higher end use pressures are required.

Another result of the study was that one can construct large single train air separation plants (up to 6000 tons/day of contained 02) rather than using multiple parallel trains. The tradeoff is between column rail shippability, overall system redundancy and the need for onsite fabrication of the large columns required. Relative costs for two air separation units using

the new process to deliver the 4250 tons/day of contained 02, required for the 500 MWe plant, are shown in table III. One unit is based on the use of three trains and rail shippable columns of 13.5 ft diameter. The other is based on a field constructed single train with a 24.3 ft column diameter. The table shows that even though the single train requires on-site fabricaton, the reduction in total number of components results in a relative capital cost savings of 11 percent.

#### Air Separation Plant Sizing

Care must be taken to determine the optimum O2 content in the oxidant, as the following calculations will show, in order to minimize the oxidant plant cost. Based on preliminary calculations, the ASU was conservatively sized for 4250 tons/day of contained oxygen in order to deliver a 35 mol percent O2 stream to the 500 MWe MHD plant. Figure 4 is a semi-logarithmic plot, which shows how the required O2 tonnage increases with O2 mol fraction in the oxidant, x. The figure also shows that the required ASU capacity is directly proportional to the "percent oxygen enrichment" parameter, PCTE (congruent curve displaced by a constant). The PCTE parameter is defined and its use is explained in the appendix.

#### IV. Calculation of 500 MW<sub>e</sub> MHD Power Plant Performance

The MHD plant performance was calculated following the methods described in previous papers.  $^{4,9}$  The power plant major cycle parameters are listed in table IV and the generator constraints are shown in table V. The bottom cycle performance was calculated using the PRESTO computer code.  $^{10}$  The bottom cycle is based on the one discussed in the AVCO CSPEC,  $^{11}$  with steam conditions of 2400 ps: $^{9}$ /1000F/1000F. The channel was cooled using low pressure, low temperature feedwater. The oxidant compressor and ASU compressors are assumed to be driven by a single steam turbine.

The generator operating conditions were selected to optimize the performance of the overall combined cycle power plant. This procedure was carried out for oxygen concentrations of 25, 30 and 35 mol percent in the oxidant, MPO, (i.e., x\*100), and channel lengths of 10 m, 15 m and 20 m.

The overall plant thermal efficiency is plotted as a function MPO for the channel lengths considered in figure 5. Results are shown (figure 5a), for an air separation plant SEC of 224 kWh/TEPO, representing the Thyssen steel plant ASU technology and also for the new process (equivalent SEC of 178 kWh/TEPO) (figure 5b). The power plant thermal efficiencies are shown to be up to 0.7 percent higher for plants using the new process (figure 5b). Also, the oxidant 02 concentration at which the maximum efficiency occurs (dashed curve) shifts to slightly higher levels. Thus, for the conventional process and a 15 m channel the maximum efficiency is about 43.7 percent, and it occurs at an MPO of 32 mol percent, (figure 5a), whereas for the new process it rises to 44.3 percent at an MPO of 33.5 mol percent, (figure 5b). Note that use of the optimized conventional air separation process

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(195 kWh/TEPO), incorporated in the ETF and the CSPEC studies, would result in overall thermal efficiency values that are about halfway between the values given above.

For both the higher and lower energy consumption, figure 5 shows that there are tradeoffs between plant thermal efficiency, oxidant 02 concentration and channel length. An additional tradeoff is shown in figure 6, which is a plot of magnetic energy stored in the channel volume as a function of plant thermal efficiency for the various exident 02 mole fractions and channel lengths considered. The magnetic energy is proportional to magnet costs while the MPO can be related to costs associated with the oxidant supply system, as indicated in figure 4. Therefore these parameters should be minimized in an optimally designed combined cycle MHD plant, consistent with sufficiently high thermal efficiency, in order to achieve the lowest cost of electricity, COE. Comparison of figures 6a and 6b shows that not only is the thermal efficiency higher at all conditions for the lower oxygen production energy, but also that a given decrease in magnetic energy will result in a smaller sacrifice in thermal efficiency for the plant with the lower oxygen production energy requirement. For example, the plant thermal efficiency is about 44.40 percent (figure 6b) using a 20 m channel and an oxidant 02 mol fraction of 0.30. For these conditions the magnetic energy is about 1150 megajoules (MJ). Reducing the channel length to 15 m, but increasing the O<sub>2</sub> mol fraction, x, to near 0.35 (actually 0.335 as per figure 5b) will result in a thermal efficiency of 44.27 percent, a loss of only 0.13 percent from the 44.4 percent figure quoted above. But the magnetic energy decreases dramatically from 1150 MJ to 675 MJ equivalent dramatically from 1150 MJ to 675 MJ, equivalent to a 41 percent decrease. Of course, the above tradeoff implies an increase in the oxidant supply system capacity of about 24 percent.

The same tradeoff, based on the results shown in figure 6a for the higher oxygen production energy requirement, will result in a power plant thermal efficiency drop from 43.90 to 43.60 percent, the resulting loss of 0.3 percent being more than double the value quoted above.

Final system optimization will require additional studies of the type discussed above, once the relationship between component sizes and associated dollar values can be defined more clearly. However, the results of this study point out that one can obtain higher overall thermal efficiencies by using large, advanced technology, oxidant supply systems which offer significant savings in oxygen production costs. These systems will also allow more flexibility in the overall power plant optimization and therefore contribute in the attainment of the lowest potential COE for combined cycle MHD/steam power plants.

#### V. Concluding Remarks

The results of this study show that the oxidant production cost for a combined cycle MHD/steam power plant can be lowered by improvements to the oxidant supply system. A new air separation process based on internal compression and liquid pumping and configured for a 500 MWe MHD/steam

power plant with a 7.22 atm oxidant delivery pressure was shown to achieve a 20 percent drop in total energy consumption below state of the art, medium purity air separation plants operating overseas. The new process also was shown to achieve a 9 percent drop below the energy requirement of optimized conventional technology air separation plants assumed for the CSPEC study. The energy consumption can be reduced further for plants requiring higher delivery pressures.

For the 500 MWe power plant, the reduced energy consumption for oxidant production was shown to result in an overall thermal efficiency increase of 0.6 percent which, when combined with projected capital cost savings identified in the study could bring about reasonable reductions in the plant COE. Future potential reductions in oxygen production costs will lead to additional improvements in MHD power plant performance.

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### TABLE I: AIR SEPARATION POWER CONSUMPTION COMPARISON FOR 500 MWe PLANT

	OPTIMIZED CONVENTIONAL PROCESS*	NEW PROCESS
INPUT AIR COMPRESSOR, KW BOOST AIR COMPRESSOR, KW EXPANSION TURBINE (POWER RECOVERY), KW PRODUCT PUMP, KW BLEND AIR COMPRESSOR, KW OXIDANT COMPRESSOR, KW	30,590 0 -620 0 0 52,990	29,630 13,410 -781 86 36,135 0
TOTAL POWER CONSUMPTION, KW	82,960	78,480

## TABLE II: AIR SEPARATION PROCESS PARAMETER COMPARISON FOR 500 MW $_{ m e}$ PLANT

	OPTIMIZED CONVENTIONAL PROCESS	NEW PROCESS
Input Air Flow, nm <sup>3</sup> /sec*	154.75	154.75
Input Air Pressure, atm	3.95	3.93
Boost Air Flow, nm <sup>3</sup> /sec		63.89
Boost Air Pressure, atm		15.99
Boost Air Temp, K	m ##	455
Upper Column Pressure, atm	1.0	1.0
Product 02 Content. mol pet.	70	70
Product Pressure, atm	1.0	7.22
Product Flow, nm <sup>3</sup> /sec	44.7	, 44.7
Product Flow, tons_02/day	4250	4250
Blend Air Flow, nm <sup>3</sup> /sec	111.3	111.3
Blend Air Pressure, atm	1.0	7.22
Blend Air Temperature, K	290	540
Oxidant Flow, nm <sup>3</sup> /sec	156.0	156.0
Oxidant O2 Content, mol pct	35	35
Oxidant Pressure, atm	7.22	7.22
Oxidant Temperature, K	544	515
SEC at Prod. Press, kWh/TEPO	195	267
SEC Corrected to P=latm, T=544K, kWh/TEPO	195	178

<sup>\*</sup> To convert nm3/sec to scfm, multiply by 5787.

## TABLE III.: COLUMN SIZE AND CAPITAL COST COMPARISONS FOR TRIPLE VS. SINGLE TRAIN ASU FOR 500 MM $_{\rm e}$ PLANT

	Triple Train	Single Train
Column Dia,* m(ft)	4.10 (13.5)	7.40 (24.3)
Relative Turnkey Cost	1.00	0.89

<sup>\*</sup>Assumed limit for rail shippability is 4270 mm (14 ft)

#### TABLE IV: MAJOR CYCLE PARAMETERS

Coal type		٠	•	٠	٠	٠		•	١	lor	Ita	m	l A	105	seb	ud
Muisture content of coal delivered to combustor, percent ,																5
Ovidizar preheat temperature K (F)	•	•	•	*	•	•	•		•	•	•	Ř	;	1	in	กั
Combiletor prelieus semperatures A (F) + + +	٠	•	*	٠	•	•	ċ.	. i.	•••		٠,		// mi	, di	i iv Imi	7/
Combustor pressure, atm	٠	٠	٠	٠	٠	٠	26	3.31	300	, Gu		,u	1116 1 & 1			46
Combustor heat loss, percent HHV of coal	•			•	•	•.		ė	þ	•			•	•	•	5
Combustor oxidizer-fuel ratio																
relative to stoichiometric Combustor slag rejection, percent	٠	•	٠	٠	•	٠	•	٠		٠	•		٠	•	u.	90
Combustor slag rejection, percent		٠	٠	٠	•	•	٠	٠	٠	٠	٠	٠	٠.			80
Generator type	٠	٠		•	٠	٠	•		•	٠	•	٠	F	ar	'ad	ay
Generator type	,	٠		٠	•	٠	•	•	٠	٠	٠	٠	٠	0.	08	59
MHD Generator inlet Mach number			•	٠									٠	٠	0	. 8
Diffuser pressure recovery coefficient .	•				٠	٠	٠	٠	٠	•	٠	,	٠		0	.6
Diffuser exit pressure, atm	٠	٠		٠		٠			٠	٠	٠	٠	٠	•	ા	.0
MHD generator length, meters	٠	•	•	,	٠	٠	٠	٠	٠	٠	•	10	),	15	5,	20
Cycle compressor polytropic efficiency .	٠				٠	٠	ě	٠	٠	٠					0.	90
Sulfur removal by seed, percent		•		•	٠			٠		•	٠	٠	•	•	•	70
Final oxidizer-fuel ratio relative																
to stoichiometric															1.	05
Stack temperature, F						•						i			2	50
Air separation plant compressor power																
requirement kW_hr/ton of equivalent																
pure oxygen added			٠										22	24.	. 1	78
Pressure drop from compressor exit to							_									
combustor percent of compressor																
exit pressure	•	•	•	٠	•	•	•	•	٠	٠	٠	٠	•	•	0	. 1
TABLE V: GENERATOR	₹ (	ON	iST	rR/	AII	NT:	S									
Manual																,
Maximum axial electric field, kV/m	٠	•	•	٠	٠	٠	٠	٠	٠	٠	•	٠	٠	•	2.	5
Maximum transverse electric field, kV/m	٠	٠	•	٠	ě	٠	●.	٠	٠		٠	٠	•	٠	4.	Ŭ
Maximum transverse current density, A/cm <sup>2</sup>	٠	٠	٠	٠	٠	٠	٠	٠		•	•	•	٠	٠	Į.	Ū
Maximum Hall parameter	٠	•		•		•	٠	٠	٠		٠	٠	٠	•	4.	Ũ
Maximum magnetic field T															6	n

#### APPENDIX: OXIDANT MIXTURE COMPOSITIONS

When air is mixed with the product delivered by medium purity air separation plants the resulting compositions, even at the same  $0_2$  mole fractions, will be different from those of pure oxygen and air mixtures. This is illustrated in tables Al and A2 which show respective compositions in terms of the mole and mass fractions of the three major constituents, N2, O2, and Ar.

The reason for the increase in concentration of argon along with that of oxygen in product streams, containing up to 95 mol percent 02, arises from the fact that the medium purity air separation process removes nitrogen, but not argon, from the product. Argon can be separated from oxygen and collected in high purity plants which use and additional argon separation column.

In MHD applications the concentration of argon in the oxidant stream has a beneficial, albeit small effect on power output, because of the slight increase in plasma conductivity. In addition to composition, tables Al and A2 also have column tabulations for mixture molecular weight, density, the ratio of contained O2 to equivalent pure O2 (which excludes any O2 contributed by air), and a parameter identified as "Percent O2 Enrichment", or PCTE. This parameter has been defined for N2-O2 mixtures 4 as

PCTE = 100 x 
$$(1 - \frac{N/0}{3.26})$$
 (1)

where N/O is the nitrogen/oxygen mass ratio in the oxidant. Equation (1) has been redefined here for multi-component mixtures as:

PCTE = 
$$100 \times (1 - \frac{R/0}{3 \times 32})$$
 (2)

where R/O is the ratio of the sum of non-oxygen mass fractions to the mass fraction of O2. Since PCTE is directly proportional to O2 plant capacity, it is a useful parameter for scaling O2 plant size when the O2 mol fraction in the oxidant is changed while, keeping the product purity constant. Tables Al and A2 show PCTE values for O2 mol fractions ranging from 21 to 100 percent for ASU product-air and pure O2-air mixtures, respectively. If the product purity is changed, the O2 plant capacity scales inversely as the ratio of the new-to-old PCTE value. Alternately, for a change in product purity, the new air separation plant capacity can be computed by multiplying by the direct ratio of "Contained O2/pure O2" values in tables Al or A2.

TABLE A1: AIR SEPARATION PLANT PRODUCT MIXTURE COMPOSITION

M Ó L OXYGEN	E PERC NITROGEN	E N T ARGON	M A S Oxygen	S PERC NITROGEN	E N T ARGON	MOLECULAR WEIGHT	DENSITY KG/M*3	CONT.02/ PURE 02	PERCENT 02 ENRICHMENT
21.00 25.00 30.00 40.00 45.00 45.00 65.00 70.00 85.00 95.00 96.00	78.064 73.885 68.663 63.440 58.217 52.994 47.771 42.548 37.325 32.102 26.879 21.653 11.211 5.988 .765	.936 1.115 1.337 1.560 1.783 2.006 2.229 2.452 2.675 2.898 3.121 3.344 3.567 3.789 4.012 4.235 4.000	23.202 27.450 32.683 37.234 42.925 52.852 57.707 62.490 67.204 71.849 76.427 80.940 85.389 89.775 94.100 95.055	75.506 71.022 65.494 60.050 54.689 49.408 44.206 39.081 34.032 29.056 24.152 19.319 14.555 9.859 5.229	1.291 1.528 1.819 2.106 2.389 2.667 2.942 3.212 3.478 3.740 3.999 4.254 4.505 4.752 4.752 4.997 5.237	28,962 29,143 29,368 29,594 29,820 30,046 30,272 30,498 30,724 30,950 31,176 31,627 31,627 31,853 32,079 32,305 32,317	1.2921 1.3002 1.3102 1.3203 1.3304 1.3405 1.3606 1.3707 1.3808 1.3909 1.4019 1.4211 1.4312 1.4413 1.4418	306.1240 4.8990 2.6325 1.9786 1.6679 1.4864 1.3673 1.2206 1.1723 1.1338 1.1024 1.0763 1.0555 1.0192 1.0159	327 20.412 37.987 50.541 59.566 67.279 73.137 77.930 81.924 85.304 88.201 90.712 92.909 94.847 96.570 98.112
97.00 98.00 99.00 100.00	.000 .000 .000	3.000 2.000 1.000 .000	96.282 97.516 98.755 100.000	.000 .000 .000	3.718 2.484 1.245 .000	32.237 32.158 32.078 31.999	1.4382 1.4347 1.4311 1.4276	1.0118 1.0077 1.0038 1.0000	98.837 99.233 99.620 100.000

TABLE A2: COMPOSITION OF PURE OXYGEN AND DRY AIR MIXTURES

M O L OXYGEN	E PER O	C E N I ARGON	M A S Oxygen	S PER NITROGEN	C E N T ARGON	MOLECULAR WEIGHT	DENSITY KG/M*3	CONT.02/ PURE 02	PERCENT 02 ENRICHMENT
21.00	78.035	.933	23.198	75.466	1.287	28.967	1.2923	327.6028	.305
25.00	74.083	.886	27.472	71.267	1.216	29.120	1.2992	4.8789	20.497
30.00	69.145	.827	32.750	66.080	1,127	29.312	1.3077	2.6203	38.164
35.00	64.206	.768	37.960	60,961	1.040	29.504	1.3163	1.9691	50.784
40.00	59.267	.709	43.102	55.908	.954	29.696	1.3248	1.6598	60.248
45.00	54.328	.650	48.179	50.920	.869	29.888	1.3334	1.4791	67.610
50.00	49.389	.591	53,190	45.996	.785	30.080	1.3420	1.3606	73.499
55.00	44.450	•532	58.138	41.134	.702	30.272	1.3505	1.2769	78.317
60.00	39.511	.473	63.024	36.333	.620	30.464	1.3591	1.2146	82,333
65,00	34.572	.414	67.849	31.592	.539	30.656	1.3677	1.1665	85.730
70.00	29.633	.354	72.613	26.911	.459	30.847	1.3762	1.1281	88.642
75.00	24.694	.295	77.319	22.287	.380	31.039	1.3848	1.0969	91.166
80,00	19.756	.236	81.966	17.720	.302	31.231	1.3933	1.0710	93.375
85.00	14.817	.177	86.557	13.209	.225	31.423	1.4019	1.0491	95.323
90.00	9.878	.118	91.093	8.752	.149	31.615	1.4105	1.0303	97.055
95.00	4.939	.059	95.573	4.350	.074	31.807	1.4190	1.0141	98.605
96.00	3.951	•047	96.463	3.476	.059	31.845	1.4207	1.0112	98.896
97.00	2.963	.035	97.350	2.604	.044	31.884	1.4225	1.0083	99.180
98.00	1.976	.024	98.236	1.734	.030	31.922	1.4242	1.0054	99.459
99.00	.988	.012	99.119	.866	.015	31.961	1.4259	1.0027	99.732
100.00	.000	.000	100.000	.000	.000	31.999	1.4276	1.0000	100.000

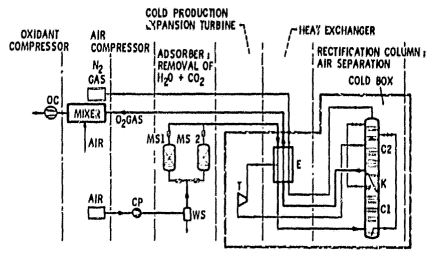


Figure 1, \* Basic air separation plant delivering product 02 at atmospheric pressure,

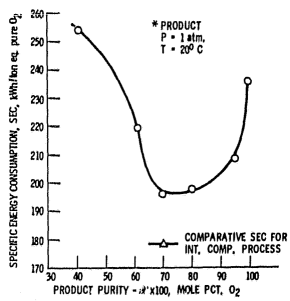


Figure 2, - Air separation process study results (Contract DEN 3-165),

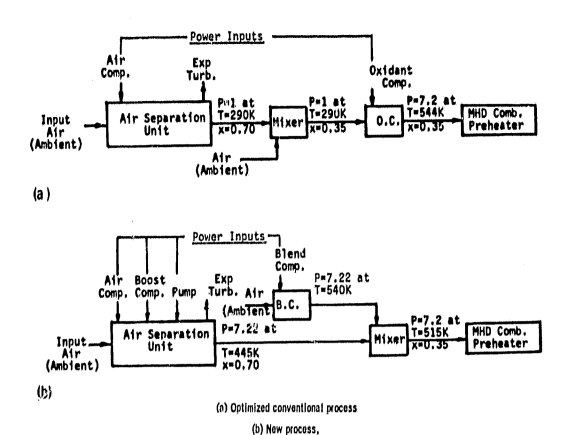


Figure 3. - Comparative block diagrams for 500 MWe MHD exidant supply systems.

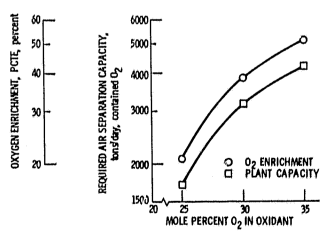
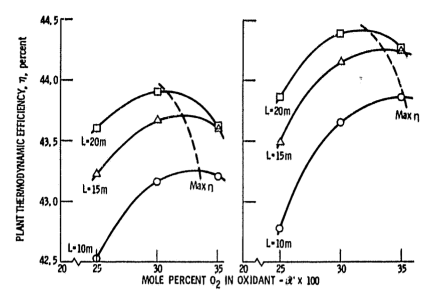
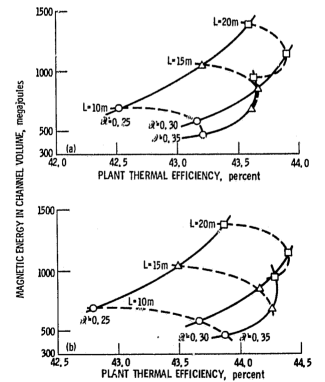


Figure 4. – Air separation plant size variation for 500 MWe  $\,$  MHD/  $\,$  steam plant. Product O  $\!_2$  concentration is 70 mole percent.



(a) Oxygen production energy is 224 kW-hr/ton of pure  $\rm O_2$ . (b) Oxygen production energy is 178 kW-hr/ton of pure  $\rm O_2$ .

Figure 5. – 500 MW  $_{\rm 0}$  MHD plant performance for various oxidant mixtures and channel lengths,



(a) Oxygen production energy is 224 kW-hr/ton of pure O<sub>2</sub>. (b) Oxygen production energy is 178 kW-hr/ton of pure O<sub>2</sub>.

Figure 6. - 500 MW<sub>e</sub> MHD plant optimization considering: magnet capacity, oxidant system capacity (α.タ), and channel length, L.

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